Air Toxic Emissions from Onroad Vehicles in MOVES3



Air Toxic Emissions from Onroad Vehicles in MOVES3

Assessment and Standards Division Office of Transportation and Air Quality U.S. Environmental Protection Agency

NOTICE

This technical report does not necessarily represent final EPA decisions or positions. It is intended to present technical analysis of issues using data that are currently available. The purpose in the release of such reports is to facilitate the exchange of technical information and to inform the public of technical developments.



Table of Contents

T	able of Co	ntents	l
L	ist of Acro	nyms	3
1	Introdu	ction	5
2	Gasolir	ne Exhaust	10
	2.1 Vo	latile Organic Compounds	10
	2.1.1	Vehicles Operating on Fuel Blends Containing 0-15 Percent Ethanol	10
	2.1.2	Vehicles Operating on Fuel Blends Containing 70-100 Percent Ethanol	40
	2.2 Po	lycyclic Aromatic Hydrocarbons (PAHs)	44
	2.2.1	Vehicles Operating on Fuel Blends Containing 0-15 Percent Ethanol	44
	2.2.2	Vehicles Operating on Fuel Blends Containing 70-100 percent Ethanol	48
	2.3 Me	etals	49
	2.4 Die	oxins and Furans	52
	2.4.1	Vehicles Operating on Fuel Blends Containing 0-15 Percent Ethanol	52
	2.4.2	Vehicles Operating on Fuel Blends containing 70-100 percent Ethanol	53
3	Diesel I	Exhaust	55
	3.1 Vo	latile Organic Compounds	55
	3.1.1	Pre-2007 Diesel Engines	55
	3.1.2	2007+ Diesel Engines	56
	3.2 Po	lycyclic Aromatic Hydrocarbons	57
	3.2.1	Pre-2007 Diesel Engines	58
	3.2.2	2007+ Diesel Engines	59
	3.3 Me	etals	60
	3.4 Die	oxins and Furans	61
4	Compre	essed Natural Gas (CNG) Exhaust	63
	4.1 Vo	latile Organic Compounds	63
	4.2 Po	lycyclic Aromatic Hydrocarbons	63
	4.3 Me	etals	64
	4.4 Die	oxins and Furans	65
5	Evapor	ative and Refueling Emissions	66
	5.1 Ga	soline Vehicles	66
	5.1.1	Vapor Venting, Fuel Leaks, and Refueling Emission Processes	66

	5.1.	2	Permeation	68
	5.2	Dies	el Vehicles	70
6	Cra	nkcas	se Emissions	71
	6.1	Vola	atile Organic Compounds	71
	6.2	Poly	cyclic Aromatic Hydrocarbons	71
	6.3	Meta	al and Dioxin Emissions	71
A	ppendi	ix A.	Development of Motor Vehicle Emission Factors for Chromium	73
A	ppendi	ix B.	Development of Motor Vehicle Emission Factors for Mercury	76
7	Ref	erenc	es	81

List of Acronyms

ACES Advanced Collaborative Emissions Study

APU auxiliary power units

CARB California Air Resource Board

CAS Chemical Abstracts Service Registry number CBD Central Business District Driving Schedule

CH4 methane

CNG compressed natural gas
CO carbon monoxide
CO2 carbon dioxide

CRC Coordinating Research Council
DOE U.S. Department of Energy
DRI Desert Research Institute
DVPE dry vapor pressure equivalent

EF emission factor

EGR exhaust-gas recirculation ETBE ethyl tertiary-butyl ether

EtOH ethyl hydroxyl

EPA U.S. Environmental Protection Agency

E0 gasoline containing 0 percent ethanol by volume
E10 gasoline containing 10 percent ethanol by volume
E15 gasoline containing 15 percent ethanol by volume
E20 gasoline containing 20 percent ethanol by volume
E70 gasoline containing 70 percent ethanol by volume
E85 gasoline containing 70-85 percent ethanol by volume

FTP Federal Test Procedure HAP hazardous air pollutants

ICP-MS inductively-coupled plasma mass spectrometry KCVES Kansas City Light-Duty Vehicle Emissions Study

LA92 Unified Driving Schedule

LIFEREG accelerated failure time regression model

LRT likelihood-ratio tests

MOBILE EPA highway vehicle emission factor model (predecessor to MOVES)

MSAT2 Control of Hazardous Air Pollutants from Mobile Sources Rule

MTBEmethyl tertiary-butyl etherNATANational Air Toxics AssessmentNEINational Emission Inventory

NERL National Exposure Research Laboratory

NVFEL EPA's National Vehicle Fuel Emissions Laboratory

NLEV National Low Emission Vehicle NMHC non-methane hydrocarbons

NMIM National Mobile Inventory Model NMOG non-methane organic gases

NOX nitrogen oxide

NREL National Renewable Energy Laboratory

OC organic carbon
Oxy oxidation catalyst

PAHs polycyclic aromatic hydrocarbons

PFI port fuel injection PM particulate matter

PUF-XAD polyurethane foam-macroreticular resins

RVP Reid Vapor Pressure

SAS Statistical Analysis System SCR selective catalytic reduction

SPECIATE repository of volatile organic gas and PM speciation profiles of air

pollution sources

SVOC semi-volatile organic compound
TAME tertiary amyl methyl ether
TBI throttle body injection

THC total hydrocarbon
TIGF teflon-impregnated glass filters

TOG total organic gases

T50 temperature at which 50 vol% of a fuel has been evaporated (°F) temperature at which 50 vol% of a fuel has been evaporated (°F)

VMT vehicle miles traveled VOC volatile organic compound

1 Introduction

The United States Environmental Protection Agency's Motor Vehicle Emission Simulator (MOVES) is a set of modeling tools for estimating air pollution emissions produced by onroad (highway) and nonroad mobile sources. MOVES estimates the emissions of greenhouse gases (GHGs), criteria pollutants and selected air toxics. The MOVES model is currently the official model for use for state implementation plan (SIP) submissions to EPA and for transportation conformity analyses outside of California. The model is also the primary modeling tool for estimating the impact of mobile source regulations on emission inventories, and thus, provides important inputs to air quality models.

Through MOVES, users can estimate inventories for selected compounds identified as air toxics in the National Emission Inventory (NEI) and National Air Toxics Assessment (NATA), and for which adequate data are available to develop mobile source emissions estimates.

The toxics included in MOVES are classified into four categories:

- 1) Volatile Organic Compounds (VOC): EPA defines VOC as any compound of carbon, excluding carbon monoxide, carbon dioxide, carbonic acid, metallic carbides or carbonates, and ammonium carbonate, which participates in atmospheric photochemical reactions, except those designated by EPA as having negligible photochemical reactivity. For use in MOVES in defining emission rates and application of speciation profiles, VOC is defined as total organic gas (TOG) minus methane, ethane, and acetone.
- 2) Polycyclic aromatic hydrocarbons (PAHs): This category is defined as hydrocarbons containing fused aromatic rings. In MOVES, the PAHs are treated separately than VOC toxics, because PAHs are semi-volatile organic compounds (SVOC) and may be measured in the gaseous phase, particulate phase, or both, depending on properties of the compound, particle characteristics and conditions in the exhaust stream or the atmosphere. MOVES estimates the sixteen PAHs (Table 1-2) that are included in the National Emissions Inventory (NEI), which does not include nitrated PAHs.
- 3) Dioxins and furans: This category includes polychlorinated organic compounds which are persistent in the environment and considered bioaccumulative in aquatic and terrestrial food chains.
- 4) Metals: This category includes metals or metal-containing compounds in elemental, gaseous and particulate phases.

Specific compounds in each category are listed in Table 1-1 through Table 1-4. Note that each compound is identified by its pollutantID in the MOVES database. With the exception of the metal species in Table 1-4, each compound is also identified by its Chemical Abstracts Service Registry number (CAS number).² For most other compounds, the identifier for the National Emissions Inventory (NEIPollutantCode in the pollutant table) is identical to the CAS number (minus the dashes). In MOVES3, methyl-tertiary-butyl-ether (MTBE) has been removed from the volatile organic compounds (Table 1-1), since it has been many years since this pollutant has been used as a gasoline additive in any significant quantity.

Table 1-1. Hydrocarbons and Volatile Organic Compounds Included in MOVES

Pollutant	pollutantID	CAS Number
Benzene	20	71-43-2
Ethanol	21	64-17-5
1,3-Butadiene	24	106-99-0
Formaldehyde	25	50-00-0
Acetaldehyde	26	75-07-0
Acrolein	27	107-02-8
2,2,4-Trimethylpentane	40	540-84-1
Ethyl Benzene	41	100-41-4
Hexane	42	110-54-3
Propionaldehyde	43	123-38-6
Styrene	44	100-42-5
Toluene	45	108-88-3
Xylene(s) ¹	46	1330-20-7

Note:

Table 1-2. Polycyclic Aromatic Hydrocarbons Included in MOVES

Pollutant	poll	utantID	CAS Number
	(gaseous phase)	(particulate phase)	
Acenaphthene	170	70	83-32-9
Acenaphthylene	171	71	208-96-8
Anthracene	172	72	120-12-7
Benz(a)anthracene	173	73	56-55-3
Benzo(a)pyrene	174	74	50-32-8
Benzo(b)fluoranthene	175	75	205-99-2
Benzo (g,h,i) perylene	176	76	191-24-2
Benzo(k)fluoranthene	177	77	207-08-9
Chrysene	178	78	218-01-9
Dibenzo(<i>a</i> , <i>h</i>)anthracene	168	68	53-70-3
Fluoranthene	169	69	206-44-0
Fluorene	181	81	86-73-7
Indeno $(1,2,3,c,d)$ pyrene	182	82	193-39-5
Naphthalene	185	23	91-20-3
Phenanthrene	183	83	85-01-8
Pyrene	184	84	129-00-0

¹ These species represent the sum of emissions from three isomers of xylene, i.e., *ortho*-, *meta*-, and *para*-xylene. In the MOVES database the sum of the different isomers of xylene is referred to simply as xylene.

Table 1-3. Dioxins and Furans Included in MOVES

Pollutant	pollutantID	CAS Number
2,3,7,8-Tetrachlorodibenzo-p-Dioxin	142	1746-01-6
1,2,3,7,8-Pentachlorodibenzo-p-Dioxin	135	40321-76-4
1,2,3,4,7,8-Hexachlorodibenzo-p-Dioxin	134	39227-28-6
1,2,3,6,7,8-Hexachlorodibenzo-p-Dioxin	141	57653-85-7
1,2,3,7,8,9-Hexachlorodibenzo-p-Dioxin	130	19408-74-3
1,2,3,4,6,7,8-Heptachlorodibenzo-p-Dioxin	132	35822-46-9
Octachlorodibenzo-p-dioxin	131	3268-87-9
2,3,7,8-Tetrachlorodibenzofuran	136	51207-31-9
1,2,3,4,6,7,8-Heptachlorodibenzofuran	144	67562-39-4
1,2,3,4,7,8,9-Heptachlorodibenzofuran	137	55673-89-7
1,2,3,4,7,8-Hexachlorodibenzofuran	145	70648-26-9
1,2,3,6,7,8-Hexachlorodibenzofuran	140	57117-44-9
1,2,3,7,8,9-Hexachlorodibenzofuran	146	72918-21-9
1,2,3,7,8-Pentachlorodibenzofuran	139	57117-41-6
2,3,4,6,7,8-Hexachlorodibenzofuran	143	60851-34-5
2,3,4,7,8-Pentachlorodibenzofuran	138	57117-31-4
Octachlorodibenzofuran	133	39001-02-0

Table 1-4. Metals Included in MOVES

Pollutant	pollutantID
Mercury (elemental gaseous)	60
Mercury (divalent gaseous)	61
Mercury (particulate)	62
Arsenic compounds	63
Chromium (Cr6+)	65
Manganese compounds	66
Nickel compounds	67

The report is structured around the four MOVES toxic categories, discussed by fuel type and emission processes. Toxics from exhaust emissions from both light-duty and heavy-duty vehicles are discussed for gasoline (Section 2), diesel (Section 3) and compressed natural gas (Section 4). For gasoline vehicles, MOVES includes the capability to estimate emissions for ethanol blends containing more than 10 percent ethanol, including E15 and E85 (70-100 percent ethanol).

The toxic emissions from the evaporative and refueling processes are discussed in Section 5, and from crankcase processes in Section 6. For exhaust, evaporative, refueling and crankcase emissions, MOVES estimates emissions from vehicles representing relevant combinations of technology and fuel; however, the availability and comprehensiveness of toxics data varies widely. Consequently, the methods and approaches used to develop model inputs also varied as necessary to incorporate the latest and best data into the model.

During MOVES runs, emissions of toxic compounds (except for metals and dioxins/furans), are estimated as fractions of the emissions of VOC, or for toxic species in the particulate phase, fractions of total organic carbon $< 2.5 \mu m$ (OC_{2.5}). MOVES estimates of VOC emissions are

calculated from emissions of total hydrocarbon (THC) as discussed in the MOVES Speciation Report.²¹ All toxic fractions are mass-based (as opposed to using molar-ratios).

For some compounds, the toxic emissions are estimated using fractions that vary with levels of fuel properties, such as ethanol, aromatics or Reid Vapor Pressure (RVP). Fractions that vary according to fuel properties are termed "complex" by MOVES. For other sets of compounds, "simple" fractions are used, meaning that the fractions are constants and do not vary with fuel properties. Note that the generalizations made here apply to evaporative as well as to exhaust emissions. In addition, in some cases, available data were sufficient to model different fractions for the two MOVES combustion processes, e.g., start and running exhaust emissions. In other cases, available data were not adequate for this purpose, with the result that the same fraction is used for both start and running emissions. Similarly, for evaporative emissions, separate fraction were developed for "permeation" and "non-permeation" processes. Finally, fractions vary with the levels of emission control (different ratios for vehicles before and after MY 2001), and for pre-2001 MY vehicles, catalyst type and fuel delivery systems.

The approach differs for estimation of emissions of metals and dioxin/furans. These species are estimated directly through application of emission rates that are assumed to be independent of operating mode. Rates for metals and dioxins/furans are expressed on a distance-specific basis (g/mile). Metals and dioxin emission rates are only produced from the 'running' exhaust emission process with the g/mile rates. Due to a lack of data, MOVES does not estimate their emissions explicitly from other exhaust emission processes such as start, extended idle, auxiliary power unit usage, and crankcase processes. However, in some cases, the start emissions for these pollutants are included in the driving cycle used to derive distance-based emission factors as discussed in the report.

The inputs used to estimate emissions of toxics do not vary with the ambient temperature simulated during a run. However, the inventories of toxic compounds estimated by the model may vary because VOC and OC_{2.5}, do vary by temperature, and as described above, emissions of toxics compounds are estimated as fractions of VOC or OC_{2.5} emissions.

Toxics inputs for MOVES are often based on very limited data, particularly for diesel and CNG engines and non-VOC pollutants. Thus, uncertainties in emission estimates may be substantial for some vehicle type/emission process/pollutant combinations. In addition, for some advanced technology vehicles, emission rates are quite low, increasing uncertainty about the toxic fraction. Additional uncertainty is introduced when we apply toxic fractions measurements from the available test programs to represent national fleet-average emissions. For VOC toxics and PAHs, this includes multiplying the fractions derived from one set of test programs by VOC emission rates from different test programs. The measurement values in this report are limited to four significant figures, to so as not to overly convey precision in the data. However, the number of significant figures should not be interpreted as the precision of the toxic emissions. Values that were not reported or were below the detection limit are listed as not-detected (ND), but are modeled as zero in MOVES. We include qualitative discussion of uncertainty, but we do not have quantitative estimates of the uncertainty of the MOVES air toxic emission estimates.

This report contains several updates made from MOVES2014b³ to MOVES3. These include minor corrections to the database to be consistent with the documentation, updated text to improve clarity of the report, and an updated speciation profile used to estimate diesel refueling toxics. Two major content updates were made to the report

First, we have removed methyl-tertiary-butyl-ether (MTBE) from the list of toxics estimated by MOVES. As discussed in the fuel supply report⁴, MOVES no longer adjusts emissions according to the gasoline oxygenate additives methyl-tertiary-butyl-ether (MTBE), ethyl-tertiary-butyl-ether (ETBE), or tertiary-amyl-methyl-ether (TAME). These oxygenates have been phased-out of the gasoline market, and we do not have data on how emissions of MY 2001 and later vehicles respond to them. For example, the EPAct fuel effects model discussed in Section 2.1.1.2 did not consider MTBE. Instead, we model historic use of MTBE in the fuel supply with matching levels of ethanol. Any equations that were functions of fuel oxygenates in MOVES2014 are now only dependent on the ethanol content of the gasoline fuel.

Second, we updated the toxic fractions and rates from MOVES2014b for model year 2007 and later diesel engines. The updated diesel exhaust emission values in Sections 3.1.2, 0, and 3.3 were peer-reviewed in September 2017. The peer-reviewed report, charge questions to the peer-reviewers, peer-review comments, EPA responses and other associated peer-review materials are located on EPA's science inventory webpage.⁵

2 Gasoline Exhaust

The gasoline toxic emissions apply to gasoline vehicles operating on gasoline with ethanol levels between 0-15 percent ethanol, as well as flex-fuel vehicles operating on high ethanol blends, with over 70% ethanol (referred to generically as E85). The toxic fractions are developed from light-duty gasoline emission rates and are applied to all gasoline vehicles, including motorcycles and heavy-duty gasoline trucks.

For volatile organic compound toxic emissions (Section 2.1), the rates are derived from two broad groups of gasoline and flex-fuel vehicles, incorporating differences in vehicle technologies, emission-control technologies and emissions standards, as well as subsets of available data and analytic methods. These two groups are defined as "model year 2000 and earlier," and "model year 2001 and later." The "model year 2001 and later" group applies to emissions starting with light-duty gasoline vehicles regulated under the National Low Emission Vehicle (NLEV) program, which began with 2001 model year vehicles, followed by the Tier 2 light-duty vehicle emission standards⁶, which began with 2004 model year vehicles and the Tier 3 light-duty vehicle emission standards, which began with 2017 model year vehicles. In addition to other technology changes due to EPA emission standards, gasoline direct injection (GDI) vehicles have entered the market; understanding the impact of GDI technology on toxics emissions is an area for future work.

For other toxic emissions from gasoline vehicles (PAHs, metals, and dioxins), we estimated fleet-average toxic emission ratios, with no distinction for vehicle technology or model year, as discussed in Sections 2.2, 2.3 and 2.4.

2.1 Volatile Organic Compounds

Within VOCs, there are separate subsections for vehicles operating on gasoline containing low ethanol percentages (Section 2.1.1) and high-ethanol percentages (Section 0).

2.1.1 Vehicles Operating on Fuel Blends Containing 0-15 Percent Ethanol

Within this sub-section, we further delineate the methods and data used for estimating toxic VOCs from 2000 and earlier vehicles (Section 2.1.1.1) and 2001 and later model year vehicles (Section 2.1.1.2).

2.1.1.1 2000 and Earlier Model Year Vehicles

Table 2-1 summarizes the methods used to estimate VOC toxic fractions. The complex model equations are stored in the MOVES tables *ComplexModelParameters*. Ethanol and acrolein are stored in the *GeneralFuelRatioExpression* table. The remaining toxic ratios based on SPECIATE profiles are stored in the *MinorHapRatio* MOVES tables. The specific data and methods used for each are described in further detail below.

Table 2-1 Calculation Methods for Gasoline Model Year 2000-and-earlier VOC Air Toxics

Compound	Fraction Type	Basis for Estimation
Benzene	complex	Complex Model
1,3-Butadiene	complex	Complex Model
Acetaldehyde	complex	Complex Model
Formaldehyde	complex	Complex Model
2,2,4-Trimethylpentane	Simple	SPECIATE profile
Acrolein	Simple	SPECIATE profile
Ethylbenzene	Simple	SPECIATE profile
n-Hexane	Simple	SPECIATE profile
Propionaldehyde	Simple	SPECIATE profile
Styrene	Simple	SPECIATE profile
Xylene(s)	Simple	SPECIATE profile
Ethanol	Simple	4 test programs outlined in Section 2.1.1.3

2.1.1.1.1 Overview of the Complex Model

For the first four compounds listed in Table 2-1, "complex" toxic fractions of VOC were estimated through application of equations developed for the Complex Model for Reformulated Gasoline. The equations are based on about 1,800 observations collected on vehicles equipped with three-way or three-way-plus-oxidation catalysts. The equations were developed by stratifying the light-duty gasoline fleet into ten technology groups (described in Table 2-2) and fitting statistical models to subsets of data for each group. The resulting sets of equations are known collectively as the "unconsolidated Complex Model." The ten groups were assigned as combinations of fuel system, catalyst type, air injection (yes/no), exhaust-gas recirculation (EGR), and normal/high emitter status. The first nine groups were intended to represent the "normal—emitting" vehicles. The tenth group represents the "high emitters," regardless of technology.

The Complex Model was designed to model the "complex" behavior of selected emissions in relation to changes in a set of selected fuel properties. The underlying dataset included measurements collected on a sample of vehicles manufactured in model year (MY) 1990 or earlier and reflecting "Tier 0" standards over a variety of gasoline formulations.

In fitting the Complex Model, the measurements for all fuel properties were "centered," meaning that the mean of all measurements for the property was subtracted from each individual measurement. This step aids in scaling the dataset so that each fuel property is centered on a mean of 0.0. Thus, if $\ln Y$ is the natural logarithm of a specific compound, such as acetaldehyde, the model is fit as shown in Equation 1 using terms for oxygenate (wt. percent), aromatics (vol. percent) and RVP (psi) as examples.

11

^a While more recent emissions data are available for Tier 1 and earlier vehicles, such as data from the Kansas City Light-Duty Vehicle Emissions Study (KCVES) mentioned later, testing was not done on a matrix of fuels which enable development of a fuel effects model.

$$ln Y = \beta_0 + \beta_{\text{oxy}} (x_{\text{oxy},i} - \bar{x}_{\text{oxy}}) + \beta_{\text{arom}} (x_{\text{arom},i} - \bar{x}_{\text{arom}}) + \cdots + \beta_{\text{RVP}} (x_{\text{RVP},i} - \bar{x}_{\text{RVP}})$$
Equation 1

The mean values used for centering all individual fuel-property values are presented in Table 2-3. Even though the input fuels in MOVES3 assume zero percent MTBE and ETBE content, because the Complex Model base fuel (see Section 2.1.1.1.2) has non-zero content of MTBE and ETBE, these centering values and coefficients are used in the Complex Model calculations. Sets of coefficients (β values in Equation 1) for models by technology group are presented for acetaldehyde, formaldehyde, benzene and 1,3-butadiene in Table 2-4 to Table 2-7. Dashes in table cells indicate no coefficient was fit for that property. It should be noted that the sulfur effects terms in the original Complex Model were not included when the model was adapted for inclusion in MOVES; rather, the sulfur effects on toxic emissions are assumed to be proportional to the effects of sulfur on total VOC, as estimated by MOVES.

Table 2-2 Technology Groups Included in the Complex Model

Technology Group	Fuel System ¹	Catalyst ²	Air Injection	Exhaust-gas Recirculation
1	PFI	3-Way	No	Yes
2	PFI	3-Way	No	No
3	TBI	3-Way	No	Yes
4	PFI	3-Way + Oxy	Yes	Yes
5	PFI	3-Way	Yes	Yes
6	TBI	3-Way	Yes	Yes
7	TBI	3-Way + Oxy	Yes	Yes
8	TBI	3-Way	No	No
9	carburetor	3-Way + Oxy	Yes	Yes
10 ("High Emitters")	ALL	ALL	ALL	ALL

Notes:

Table 2-3 Mean Fuel-Property Values Used for Centering Terms in the Complex Model

Property	Units	Mean Value
Aromatics	Vol. %	28.26
Olefins	Vol. %	7.32
Methyl-tertiary-butyl-ether (MTBE) ¹	Wt.%	0.95
Ethyl-tertiary-butyl-ether (ETBE) ¹	Wt. %	0.023
Ethanol (EtOH) ¹	Wt. %	0.314
Tertiary-amyl-methyl-ether (TAME) ¹	Wt. %	0.016
Oxygenate ²	Wt. %	1.77
RVP	Psi	8.61
E200	%	46.73
E300	%	85.90

Notes:

¹ Fuel System: PFI = port fuel injection, TBI = throttle body injection.

² Catalyst: "3-way" = three-way catalyst, "Oxy" = oxidation catalyst.

¹ Species-specific values used in the aldehyde models.

² Aggregate value used for the butadiene and benzene models. As calculated using the volumes of MTBE, ETBE, EtOH, and TAME and additional parameters documented in Appendix B of the MOVES Fuel Effects Report²²

Table 2-4 Complex Model Coefficients for Acetaldehyde, by Technology Group

Fuel Property									
Technology Group	Aromatics	Olefins	\mathbf{MTBE}^1	\mathbf{ETBE}^1	ЕŧОН	TAME	RVP	E200	E300
1	-0.055	-	-0.036	0.316	0.249	-	-	-	-0.012
2	-0.055	-	-	0.316	0.249	-	-	-	-0.012
3	-0.055	-	-	0.316	0.249	-	-	-	-0.012
4	-0.055	-	-	0.316	0.249	-	0.242	-	-0.012
5	-0.055	-	-	0.316	0.249	-	1	-	-0.012
6	-0.055	-	-	0.316	0.249	-	-	-	-0.012
7	-0.055	-	-	0.316	0.249	-	-	-	-0.012
8	-0.055	-	-	0.316	0.249	-	-	_	-0.012
9	-0.055	-	-	0.316	0.249	-	-	_	-0.012
10	-0.055	-	-0.056	0.316	0.249	-	-	-	-0.012

Notes:

Table 2-5 Complex Model Coefficients for Formaldehyde, by Technology Group

					el Propei		Technology Group					
Technology Group	Aromatics	Olefins	MTBE ¹	ETBE	ЕтОН	TAME	RVP	E200	E300			
1	-0.007	1	0.046	1	Ī	1	1	-	-0.010			
2	-0.007	1	0.046	1	Ī	1	1	-	-0.010			
3	-0.007	ı	0.046	ı	Ī	ı	1	-	-0.010			
4	-0.007	ı	0.046	ı	Ī	ı	1	-	-0.010			
5	-0.007	ı	0.046	ı	Ī	ı	1	-	-0.010			
6	-0.007	ı	0.046	ı	Ī	ı	1	-	-0.010			
7	-0.007	ı	0.046	ı	Ī	ı	1	-	-0.010			
8	-0.007	-	0.046	-	-	-	-	-	-0.010			
9	-0.007	-	0.046	-	-	-	-	-	-0.010			
10	-0.007	-0.031	0.046	-	-	-	-	-	-0.010			

Notes:

¹ Even though the input fuels in MOVES3 assume 0 percent MTBE and ETBE content, because the base fuel has non-zero content of MTBE and ETBE, these coefficients are used in the Complex Model calculations (Table 2-3)

¹ Even though the input fuels in MOVES3 assume 0 percent MTBE content, because the base fuel has non-zero content of MTBE, these coefficients are used in the Complex Model calculations (Table 2-3)

Table 2-6 Complex Model Coefficients for Exhaust Benzene, by Technology Group

			F	uel Propert	ty	8,	-
Technology Group	Aromatics	Olefins	Oxygenate	Fuel Benzene	RVP	E200	E300
1	0.026	-	-	0.222	-	-0.0095	-
2	0.026	-	-	0.222	-	-	-
3	0.026	-	-	0.222	-	-0.0058	-
4	0.026	-	ı	0.222	-	ı	-
5	0.049	-	ı	0.222	-	ı	-
6	0.026	-	ı	0.222	-	ı	-
7	0.026	-	ı	0.222	-	ı	-
8		-	-	0.222	-	-	-
9	0.026	-	-	0.222	-	-	-
10	0.012	-	-0.096	0.222	-	-	0.011

Table 2-7 Complex Model Coefficients for 1,3-Butadiene, by Technology Group

	Fuel Prope						
Technology Group	Aromatics	Olefins	Oxygenate	E200	E300		
1	-0.004	0.028	-	-0.0073	-0.017		
2	-0.004	0.028	-	-0.0073	-0.017		
3	-0.004	0.028	-	-0.0073	-0.006		
4	-0.004	0.028	-	-0.0073	-0.017		
5	-0.004	0.028	-	-0.0073	-0.017		
6	-0.004	0.028	-	0.0058	-0.017		
7	-0.004	0.028	-	-0.0073	-0.017		
8	-0.004	0.028	-	-0.0073	-0.017		
9	-0.004	0.028	-	-0.0073	-0.017		
10	-0.004	0.044	-0.061	-0.0073	-0.008		

2.1.1.1.2 Application of the Complex Model

In MOVES, the complex model equations are consolidated by weighting them together using model-year specific weights based on the mix of technologies in the sales fleet for each model year, as obtained from MOBILE6.2.⁵⁴

For each compound, Equation 1 is used to estimate the effects of both "base" and "target" fuels. We assume that vehicles were running on a specific fuel when the data underlying the base emission rates were measured. We refer to these fuels as "base" fuels and use them as reference points to estimate the effects of "target" fuels simulated during MOVES runs.²² The "target" fuels are represented by specific sets of fuel properties and represent fuels "in-use" in the geographic area(s) and season(s) being modeled in MOVES.

Initially, an adjustment for the difference in emissions of the compound modeled on the target fuel relative to the base fuel is calculated. If the model, as shown in Equation 2, is expressed, using matrix notation as $X\beta_{target}$ and $X\beta_{base}$ for estimates on the target and base fuels, then the fractional difference in emissions is given by:

$$f_{\text{adj}} = \frac{exp(X\beta_{\text{target}})}{exp(X\beta_{\text{base}})} - 1.0$$
 Equation 2

The expression in Equation 2 is evaluated for target and base fuels for each of the ten technology groups. A mean value of the adjustment is then calculated for each model year from 2000 back to 1970, as a weighted average of the fraction of sales in each group in each model year, for the groups, as shown in Equation 3. The weights are shown in Table 2-8. The weights represent the sales fractions and high emitter fractions for the ten vehicle technologies defined in Table 2-2 above.

Note that the use of varying weights in applying the Complex Model in MOVES differs from the original application in which the weights were invariant. The application of Equation 3 to each of the thirty ages listed in Table 2-8 gives a set of 30 adjustments; the one that is applied for each model year depends on the calendar year simulated.

$$f_{
m adj,mean} = \sum_{Group=1}^{10} w_{
m Group} f_{
m adj,Group} \quad ; \quad \sum_{Group=1}^{10} w_{Group} = 1.0$$
 Equation 3

The mean adjustments calculated in Equation 3 are then applied to estimate emissions of the toxic on the target fuel ($E_{\rm relative,toxic}$), representing the effect on the emissions of the toxic due to the changes in fuel properties between the target and base fuels. If the target and base fuels were identical, the values of $f_{\rm adj,mean}$ would be 0.0.

$$E_{\text{relative,toxic}} = E_{\text{base,toxic}} (1 + f_{\text{adi,mean}})$$
 Equation 4

The calculations in Equation 1 to Equation 4 are also applied to VOC emissions, ending with the generation of a value of $E_{\text{relative,VOC}}$. The Complex Model coefficients for VOC are documented in the rulemaking analysis.⁸ This value for VOC is then combined with that of each toxic to calculate a fraction of VOC used to estimate the total mass of emissions for each toxic during a model run. These fractions are denoted as f_{toxic} and calculated as shown in Equation 5.

$$f_{\text{toxic}} = \frac{E_{\text{relative,toxic}}}{E_{\text{relative,VOC}}}$$
 Equation 5

As a final step, the mass emissions of each toxic (I_{toxic}) during a model run are estimated by multiplying the mass of VOC emissions estimated by MOVES (I_{VOC}) by the values of f_{toxic} .

$$I_{\text{toxic}} = f_{\text{toxic}} I_{\text{VOC}}$$
 Equation 6

The equations and parameters presented are used to estimate the fuel impacts for both Tier 0 and Tier 1 gasoline vehicles. This approach is based on the assumption that the proportional responses of air toxic emissions to changes in fuel properties are similar for vehicles certified to both sets of standards.

The Complex Model equations are applied to running, start and extended idle emissions for gasoline-fueled vehicles for all 2000 and earlier model years for the first four pollutants listed in Table 2-1 (acetaldehyde, formaldehyde, benzene and 1,3-butadiene). In addition, MOVES applies the Complex Model based on light-duty gasoline vehicles to heavy-duty gasoline vehicles. This step was taken because the very limited data specific to heavy-duty gasoline vehicles used in MOBILE6.2 were considered not adequate to accurately capture the effects of fuel properties on those vehicles.

Table 2-8 Weights Applied to Complex Model Coefficients for Technology Groups, by Age (Vehicle Age 0 Represents Model Year 2000)^b

Age					Technolo					
	1	2	3	4	5	6	7	8	9	10
0	0.2360	0.2829	0.1806	0.1814	0.0290	0.0042	0.0556	0.0	0.0203	0.0100
1	0.2339	0.2803	0.1789	0.1797	0.0287	0.0042	0.0551	0.0	0.0201	0.0190
2	0.2315	0.2774	0.1771	0.1779	0.0284	0.0041	0.0546	0.0	0.0199	0.0290
3	0.2272	0.2723	0.1738	0.1746	0.0279	0.0041	0.0536	0.0	0.0196	0.0470
4	0.2229	0.2672	0.1706	0.1713	0.0274	0.0040	0.0525	0.0	0.0192	0.0650
5	0.2189	0.2623	0.1675	0.1682	0.0269	0.0039	0.0516	0.0	0.0188	0.0820
6	0.2148	0.2574	0.1644	0.1651	0.0264	0.0038	0.0506	0.0	0.0185	0.0990
7	0.2110	0.2529	0.1614	0.1621	0.0259	0.0038	0.0497	0.0	0.0182	0.1150
8	0.2072	0.2483	0.1585	0.1592	0.0254	0.0037	0.0488	0.0	0.0178	0.1310
9	0.2036	0.2440	0.1558	0.1565	0.0250	0.0036	0.0480	0.0	0.0175	0.1460
10	0.2000	0.2397	0.1530	0.1537	0.0246	0.0036	0.0471	0.0	0.0172	0.1610
11	0.1967	0.2357	0.1505	0.1512	0.0241	0.0035	0.0464	0.0	0.0169	0.1750
12	0.1934	0.2317	0.1479	0.1486	0.0237	0.0035	0.0456	0.0	0.0166	0.1890
13	0.1903	0.2280	0.1456	0.1462	0.0234	0.0034	0.0448	0.0	0.0164	0.2020
14	0.1872	0.2243	0.1432	0.1438	0.0230	0.0033	0.0441	0.0	0.0161	0.2150
15	0.1843	0.2209	0.1410	0.1416	0.0226	0.0033	0.0434	0.0	0.0159	0.2270
16	0.1814	0.2174	0.1388	0.1394	0.0223	0.0032	0.0428	0.0	0.0156	0.2390
17	0.1786	0.2140	0.1366	0.1372	0.0219	0.0032	0.0421	0.0	0.0154	0.2510
18	0.1760	0.2109	0.1346	0.1352	0.0216	0.0031	0.0415	0.0	0.0151	0.2620
19	0.1736	0.2080	0.1328	0.1334	0.0213	0.0031	0.0409	0.0	0.0149	0.2720
20	0.1712	0.2052	0.1310	0.1315	0.0210	0.0031	0.0403	0.0	0.0147	0.2820
21	0.1688	0.2023	0.1291	0.1297	0.0207	0.0030	0.0398	0.0	0.0145	0.2920
22	0.1664	0.1994	0.1273	0.1279	0.0204	0.0030	0.0392	0.0	0.0143	0.3020
23	0.1643	0.1969	0.1257	0.1262	0.0202	0.0029	0.0387	0.0	0.0141	0.3110
24	0.1624	0.1946	0.1242	0.1248	0.0199	0.0029	0.0383	0.0	0.0140	0.3190
25	0.1602	0.1920	0.1226	0.1231	0.0197	0.0029	0.0378	0.0	0.0138	0.3280
26	0.1602	0.1920	0.1226	0.1231	0.0197	0.0029	0.0378	0.0	0.0138	0.3280
27	0.1602	0.1920	0.1226	0.1231	0.0197	0.0029	0.0378	0.0	0.0138	0.3280
28	0.1602	0.1920	0.1226	0.1231	0.0197	0.0029	0.0378	0.0	0.0138	0.3280
29	0.1602	0.1920	0.1226	0.1231	0.0197	0.0029	0.0378	0.0	0.0138	0.3280
30	0.1602	0.1920	0.1226	0.1231	0.0197	0.0029	0.0378	0.0	0.0138	0.3280

2.1.1.1.3 Simple Fractions of VOC

MOVES uses a simpler approach to model emissions of 2,2,4-trimethylpentane, acrolein, ethylbenzene, n-hexane, propionaldehyde, styrene, xylene(s) and ethanol. Table 2-9 lists toxic fractions of VOC for these compounds for a range of gasoline-ethanol blends. With the exception of ethanol, these fractions were developed for E0 and E10 by Sierra Research using speciation profiles estimated from EPA's SPECIATE 4.2 database.⁹

_

^b Note that in the MOVES database, these weights are stored in the table FuelModelWtFactor.

Emissions of ethanol in exhaust are estimated for gasoline blends containing ethanol at levels of 0 to 10 vol percent. For vehicles running on 10 percent ethanol, ethanol was estimated to comprise 2.39 percent of exhaust VOC. This estimate is based on results measured on nine vehicles in four test programs. ^{10,11,12,13} The fraction of ethanol in exhaust VOC for blends containing 5.0 percent and 8.0 percent ethanol is estimated by interpolating linearly between the fractions for 0.0 percent and 10.0 percent ethanol.

No data exist for MY 2000 and earlier vehicles running on E15 or E20. Pre-2000 MY gasoline vehicles do not have an EPA waiver to operate on ethanol fractions higher than 10 percent. ^{14,c} For acrolein and ethanol, we simply extended the E10 toxic fractions as shown in Table 2-9. For pollutantIDs 40 – 46, we used toxics ratios for 2001 and later vehicles for E15 and E20, found in Table 2-35.

Table 2-9 Toxic Fractions of VOC for Selected Air Toxics, Representing Gasoline and Ethanol Blends

Compound	pollutantID	Fuel Blend (by Ethanol Level)					
-		0% (E0)	10% (E10)	15% (E15)	20% (E20)*		
Ethanol	21	0	0.024	0.024	0.024		
Acrolein	27	0.00063	0.00063	0.00063	0.00063		
2,2,4- Trimethylpentane	40	0.018	0.018	0.022	0.0046		
Ethyl Benzene	41	0.021	0.019	0.016	0.022		
Hexane	42	0.016	0.016	0.011	0.025		
Propionaldehyde	43	0.00086	0.00086	0.00060	0.00066		
Styrene	44	0.0011	0.0010	0.0046	0.0041		
Toluene	45	0.096	0.087	0.073	0.096		
Xylene	46	0.078	0.070	0.069	0.093		

^{*}Note, MOVES versions after MOVES2014a do not estimate emissions for vehicles using E20 fuel, but these fractions currently remain in the MOVES database.

In the MOVES database, these inputs are stored in the table "minorHAPratio." In the label, the term "HAP" refers to "hazardous air pollutant." In MOVES, we refer to these pollutants stored in this table as minor HAPs. A description of the table is provided in Table 2-10.

use cases, in which MOVES would use the E15 toxic fractions in Table 2-9. In MOVES, all model year gasoline vehicles use the same market of gasoline fuels specified in the fuel supply table. Ideally, MOVES would not model E15 usage from MY 2000 and earlier vehicles when users specify E15 marketshare for their runs. However, we have not incorporated this additional complexity into the model due to the small contribution of MY 2000 vehicles to current emission inventories from E15 fuel blends.

18

^c Due to the small volume of E15 fuel sales and the uncertainty of allocating the E15 across the country, MOVES3 assumes no marketshare of E15 in the default fuel supply. Users can specify marketshare of E15 for the individual

Table 2-10 Description of the Database Table "minorHAPRatio"

	e 2-10 Description of the Database 18	I
Field	Description	RelevantValues
polProcessID	Identifies the pollutant (1st two digits and Emissions Process (last two digits).	Pollutants are identified in the table above; Relevant processes include: "Running Exhaust" (processID = 1) "Start Exhaust" (processID = 2)
fuelTypeID	Identifies broad classes of fuels, e.g., "gasoline." "diesel."	1 = "Gasoline" 2 = "Diesel" 5 = "Ethanol"
fuelSubTypeID	Identifies specific fuel classes within the fuelTypeID	10 = "Conventional Gasoline" 11 = "Reformulated Gasoline" 12 = "Gasohol (E10)" 13 = "Gasohol (E8)" 14 = "Gasohol (E5)" 15 = "Gasohol (E15)" 18 = "Gasohol (E20)"* 51 = "Ethanol (E85)" 52 = "Ethanol (E70)"
modelYearGroupID	Identifies a set of model years covered by a specific value of atRatio.	1960-1970 1971-1977 1978-1995 1996-2003 2004-2060
atRatio	Fraction, or "ratio" of the toxic relative to total VOC.	
atRatioCV	"Coefficient of Variation of the Mean" or "relative standard error" of the atRatio.	
dataSourceID	Indicates source data and methods used to estimate atRatio.	

^{*}Note MOVES versions after MOVES2014a do not estimate emissions for vehicles using E20 fuel.

2.1.1.2 2001 and later model year vehicles

For vehicles manufactured in model year 2001 and later and certified to NLEV or Tier 2 standards more recent data were available. As for the earlier model years, toxic emissions are estimated as fractions of VOC, with toxic fractions for various compounds estimated using differing datasets and methods. For some compounds and processes, models were developed to estimate "complex" fractions (as a function of gasoline properties), whereas for others, "simple" constant fractions were estimated. In some cases, different fractions could be estimated for the start and running emission processes. For the compounds included in MOVES, data sources and estimation methods are summarized in Table 2-11. The data used to estimate the toxic fractions were obtained from the EPAct Program which is discussed in the following section. For benzene and 1,3-butadiene there are additional adjustments based on the benzene and olefin content of the fuel as discussed in Section 2.1.1.2.3.

Table 2-11 Data Sources and Methods Used to Estimate Gasoline Model Year 2001-and later Toxic Fractions for VOCs

Compound	Process	Fraction Type	Basis for Estimation
Acetaldehyde	Start	complex	application of EPAct models ¹
	Running	complex	application of EPAct models
Formaldehyde	Start	complex	application of EPAct models
	Running	complex	application of EPAct models
Acrolein	Start	complex	application of EPAct models
	Running	simple	Data from EPAct Project (Phase 3) ²
Ethanol	Start	complex	application of EPAct models
	Running	complex	application of EPAct models
Benzene*	Start	complex	application of EPAct models
	Running	simple	Data from EPAct Project (Phase 3)
1,3-Butadiene*	Start	complex	application of EPAct models
	Running	simple	Data from EPAct Project (Phase 3)
2,2,4-Trimethylpentane	Both	simple	Speciation Profile (EPAct Phase 1) ³
Ethylbenzene	Both	simple	Speciation Profile (EPAct Phase 1)
N-Hexane	Both	simple	Speciation Profile (EPAct Phase 1)
Propionaldehyde	Both	simple	Speciation Profile (EPAct Phase 1)
Styrene	Both	simple	Speciation Profile (EPAct Phase 1)
Xylene(s)	Both	simple	Speciation Profile (EPAct Phase 1)

^{*}Additional adjustments are made to the benzene and 1,3-butadiene emissions as discussed in Section 2.1.1.2.3.

2.1.1.2.1 Overview of the EPAct Program

To better understand the effect of gasoline fuel properties on exhaust emissions from Tier-2 certified vehicles, EPA entered a partnership with the Department of Energy (DOE) and the Coordinating Research Council (CRC). The resulting research program was dubbed the "EPAct/V2/E-89" program (or "EPAct" for short). The program was conducted in three phases. Phases 1 and 2 were pilot efforts involving measurements on 19 light-duty cars and trucks using three fuels, at two temperatures. These preliminary efforts laid the groundwork for design of a full-scale research program, designated as Phase 3.

Initiated in March 2009, the Phase 3 program involved measurement of exhaust emissions from fifteen high-sales-volume Tier-2 certified vehicles. The vehicles were selected to represent the latest technologies in the market at the time the program was launched (2008). The vehicles were to reflect a majority of sales for model year 2008. In addition, the vehicles were to conform primarily to Tier-2 Bin-5 exhaust standards, and to reflect a variety of emission-control technologies, as realized through the selection of a range of vehicle sizes and manufacturers. The vehicle sample is summarized in Table 2-12.

Table 2-12 Test Vehicles for the Phase-3 EPAct Program (all vehicles in MY2008)

Make	Brand	Model	Engine Size	Tier 2 Bin	LEVII	Odometer
					Std	
GM	Chevrolet	Cobalt	2.2L I4	5	NA	4,841
GM	Chevrolet	Impala FFV	3.5L V6	5	L2	5,048
GM	Saturn	Outlook	3.6L V6	5	L2	5,212
GM	Chevrolet	Silverado FFV	5.3L V8	5	NA	5,347
Toyota	Toyota	Corolla	1.8L I4	5	U2	5,019
Toyota	Toyota	Camry	2.4L I4	5	U2	4,974
Toyota	Toyota	Sienna	3.5L V6	5	U2	4,997
Ford	Ford	Focus	2.0L I4	4	U2	5,150
Ford	Ford	Explorer	4.0L V6	4	NA	6,799
Ford	Ford	F150 FFV	5.4L V8	8	NA	5,523
Chrysler	Dodge	Caliber	2.4L I4	5	NA	4,959
Chrysler	Jeep	Liberty	3.7L V6	5	NA	4,785
Honda	Honda	Civic	1.8L I4	5	U2	4,765
Honda	Honda	Odyssey	3.5L V6	5	U2	4,850
Nissan	Nissan	Altima	2.5L I4	5	L2	5,211

The study used a total of twenty-seven test fuels spanning wide ranges of five fuel properties (ethanol, aromatics, vapor pressure, and two distillation parameters: T50 and T90). The number of test points and values of each property are shown in Table 2-13. The properties of the test fuels were not assigned to represent in-use fuels, but rather to allow development of statistical models that would enable estimation of relative differences in emissions across the ranges of fuel properties expected in commercially available summer fuels in the U.S. (5th to 95th percentiles for each property).

Table 2-13 Levels Assigned to Experimental Factors (Fuel parameters) for the Phase-3 EPAct Program

Factor	No. Levels	Levels				
		Low	Middle	High		
Ethanol (vol.%)	4	0	10, 15	20		
Aromatics (vol.%)	2	15		35		
RVP (psi)	2	7		10		
T50 (°F)	5	150	165, 190, 220	240		
T90 (°F)	3	300		340		

The LA92 test cycle was used, with emissions measured over three phases analogous to those in the Federal Test Procedure (FTP), at an ambient temperature of 75°F. Note that throughout this chapter, the terms "start," "cold start" and "Bag 1" will be treated as synonymous, and similarly, the terms "running," "hot-running" and "Bag 2" will also be treated as synonymous.

The experimental design embodied in the fuel set is the product of an iterative process involving balancing among research goals, fuel-blending feasibility and experimental design. As fuel properties tend to be moderately to strongly correlated, and as the goal was to enable analysis of fuel effects as though the properties were independent (uncorrelated), it was necessary to address these issues in design and analysis. Accordingly, the fuel set was designed using a computergenerated optimal design, as modified by additional requirements such as the total number of fuels and specific properties for subsets of fuels. In addition, to generate the design, it was

necessary to specify the fuel effects to be estimated by the resulting model. The fuel set was designed to allow estimation of linear effects for the five properties shown in Table 2-13, plus two-way interactions of ethanol and the other five properties, as shown in Equation 7, in which β represents a linear coefficient for each effect.

$$Y = \beta_0 + \beta_1 \text{etOH} + \beta_2 \text{Arom} + \beta_3 \text{RVP} + \beta_4 T 50 + \beta_5 T 90 + \beta_6 T 50^2 + \beta_{11} \text{etOH}^2$$

$$\beta_7 \text{etOH} \times \text{Arom} + \beta_8 \text{etOH} \times \text{RVP} + \beta_9 \text{etOH} \times \text{T50} + \beta_{10} \text{etOH} \times \text{T90}$$

$$+ \varepsilon$$
Equation 7

In the equation, the linear terms (e.g., β_1 etOH, etc.) describe linear associations between emissions (Y) and the value of the fuel property. The quadratic terms are used to describe some degree of curvature in the relationship between emissions and the fuel property. Note that a minimum of three test levels for a property is needed to assess curvilinear relationships and that the design included such effects only for ethanol and T50. Two-way interaction terms indicate that the relationship between emissions and the first fuel property is dependent on the level of the second fuel property. For example, if an etOH×Arom interaction is included in a model, it implies that the effect of ethanol on the emission Y cannot be estimated without accounting for the aromatics level, and vice versa. Note that inclusion of the 11 effects in the design does not imply that all effects will be retained in all models following the fitting process. Properties for each of the test fuels are shown in Table 2-14.

Emissions measured include carbon dioxide (CO₂), carbon monoxide (CO), THC, methane (CH₄), oxides of nitrogen (NO_X), and PM_{2.5}. In addition, hydrocarbons were speciated for subsets of vehicles and fuels, allowing calculation of derived parameters such as non-methane organic gases (NMOG) and non-methane hydrocarbons (NMHC). Speciation also allowed independent analyses of selected toxics including acetaldehyde, formaldehyde, acrolein, benzene, 1,3-butadiene and ethanol.

Due to limitations in budget, the entire study design was not applied to speciated hydrocarbons, including those discussed in this chapter. For the speciated compounds, the number of relevant measurements varies by bag, compound and vehicle. For selected compounds, measurements for Bag 1 were taken for all vehicles over the entire fuel set, thus encompassing the entire study as designed, including replication. However, for the remaining compounds in Bag 1 and for all compounds in Bags 2, measurements were taken for a smaller number of vehicles over a reduced set of fuels, without replication. The combinations of fuels and vehicles included for each compound analyzed are summarized in Table 2-15.

Throughout this chapter, the complete set of 27 fuels will be denoted as the "full design," as it includes all the fuel parameter points for which the design was optimized. Similarly, the set of 11 fuels will be denoted as the "reduced design," as it covers a set of fuel parameter points narrower than that for which the design was originally optimized. Note that Table 2-14 also identifies the subset of fuels included in the reduced design.

Phase 3 data collection was completed in June 2010. Dataset construction and analysis was conducted between January 2010 and November 2012. This process involved ongoing

collaboration among EPA staff, Department of Energy (DOE) staff and contractors, and CRC representatives. Following the completion of data collection, construction of the dataset involved intensive evaluation and quality assurance. The analysis involved several iterations between analysis and additional physical and chemical review of the data. Successive rounds of statistical modeling were applied to the data to achieve several goals, including identification of potential candidate models, identification and review of outlying observations, identification and review of subsets of data from influential vehicles, and identification of models including subsets of terms that best explain the results obtained. The EPAct exhaust research program and analysis are extensively documented in the "EPAct Test Program Report". In an "EPAct Analysis Report". In an analysis are extensively documented in the "EPAct Test Program Report".

This document describes how the data and statistical models developed during the EPAct study are applied to model toxics in the MOVES model.

Table 2-14 Measured Parameters for Fuels in the Phase-3 EPAct Program

I at	Table 2-14 Measured Parameters for Fuels in the Phase-3 EPAct Program								
Fuel ¹	etOH (vol.%)	Aromatics (vol.%)	RVP (psi)2	T50 (°F)	T90 (°F)				
1	10.03	15.4	10.07	148.9	300.2				
2	0	14.1	10.2	236.7	340.1				
3^{3}	10.36	15.0	6.93	217.5	295.9				
4	9.94	15.5	10.01	221.9	337.5				
5	0	34.7	6.95	237.0	300.0				
6^{3}	10.56	15.0	7.24	188.5	340.4				
7^{3}	0	17.0	7.15	193.1	298.4				
8	0	15.7	10.2	221.1	303.1				
9	0	35.8	10.30	192.8	341.8				
10^{3}	9.82	34.0	7.11	217.1	340.2				
11	10.30	35.0	9.93	189.3	298.6				
12	9.83	34.8	10.13	152.2	339.8				
13^{3}	0	34.1	6.92	222.5	337.9				
143	0	16.9	7.14	192.8	338.5				
15	0	35.3	10.23	189.7	299.4				
16	10.76	35.6	7.12	218.8	300.6				
20	20.31	15.2	6.70	162.7	298.7				
21^{3}	21.14	35.5	7.06	167.6	305.0				
22	20.51	15.0	10.21	163.2	297.3				
23^{3}	20.32	15.9	6.84	162.5	338.2				
24	20.51	15.3	10.12	165.1	338.1				
25	20.03	35.2	10.16	166.9	337.9				
26	15.24	35.6	10.21	160.3	338.7				
273	14.91	14.9	6.97	221.5	340.3				
28^{3}	14.98	34.5	6.87	216.6	298.8				
30	9.81	35.5	10.23	152.9	323.8				
31^{3}	20.11	35.5	6.98	167.3	325.2				

Notes:

¹ Note that numbering of fuels is not entirely sequential throughout.

² This parameter was measured as "DVPE," but for simplicity, will be referred to as "RVP" in this document.

³ These fuels included in the "reduced design."

Table 2-15 Features of the Study Design Applied to Speciated Compounds Selected for Analysis

Compound	Bag 1			Bag 2		
	No. vehicles	No. Fuels	replication	No. vehicles	No. Fuels	replication
Acetaldehyde	15	27	YES	5	11	NO
Formaldehyde	15	27	YES	5	11	NO
Acrolein	15	27	YES	5	11	NO
Ethanol	15	27	YES	5	11	NO
Benzene	15	11	NO	5	11	NO
1,3-Butadiene	15	11	NO	5	11	NO
Ethane	15	11	NO	5	11	NO

Standardizing Fuel Properties

In model fitting, as well as in applying the resulting sets of coefficients, it is necessary to first "center" and "scale" the properties of fuels, also known as "standardization." This process simply involves first "centering" the measured fuel properties by subtracting the sample mean from the given value, and then "scaling" by then dividing the centered values by their respective standard deviations, as shown in Equation 8. Note that the means and standard deviations are calculated from the fuel set used for the program (see Table 2-14). The result is a "Z score," representing a "standard normal distribution" with a mean of 0.0 and a standard deviation of 1.0.

$$Z_i = \frac{x_i - \bar{x}}{S}$$
 Equation 8

For the linear effects in the model, standardization is performed using the values of each fuel property, each in their respective scales (vol. percent, psi, °F.). Using aromatics as an example, the standardization of the linear term is shown in Equation 9.

$$Z_{\text{arom}} = \frac{x_{\text{arom}} - \bar{x}_{\text{arom}}}{S_{\text{arom}}}$$
 Equation 9

For second-order terms, however, the process is not performed on the values of the fuel properties themselves. Rather, quadratic and interaction terms are constructed from the Z scores for the linear terms, and the process is repeated. Using the quadratic term for ethanol as an example (etOH×etOH), the standardized value, denoted by $ZZ_{\text{etOH}\times\text{etOH}}$, is calculated as shown in Equation 10, where $m_{Z_{\text{etoH}}Z_{\text{etoH}}}$ and $s_{Z_{\text{etoH}}Z_{\text{etoH}}}$ are the mean and standard deviation of the quadratic term constructed from the Z score for the linear effect.

$$ZZ_{\text{etOH} \times \text{etOH}} = \frac{Z_{\text{etOH}} Z_{\text{etOH}} - m_{Z_{\text{etOH}} Z_{\text{etOH}}}}{s_{Z_{\text{etOH}} Z_{\text{etOH}}}}$$
 Equation 10

Standardized terms for interaction effects are constructed similarly. For example, Equation 11 shows the standardization of an interaction term between ethanol and aromatics.

$$ZZ_{\text{etOH} \times \text{eArom}} = \frac{Z_{\text{etOH}} Z_{\text{Arom}} - m_{Z_{\text{etOH}} Z_{\text{Arom}}}}{S_{Z_{\text{etOH}} Z_{\text{Arom}}}}$$
 Equation 11

Means and standard deviations for relevant model terms are shown in Table 2-16. Note that the means and standard deviations shown in the table are calculated from the fuel set itself as shown in the table; in this calculation the properties are not weighted for numbers of replicates on each fuel and emission combination. In this way, the process is simplified by using the same standardization in fitting all models, as well as in subsequent applications of the models. Note also that the reduced fuel set is standardized using a different set of parameters than the full fuel set.

The process of standardization is illustrated for three test fuels in Table 2-17. Overall, the process applied here is similar to the "correlation transformation" sometimes applied in multiple regression. One difference in this case is that the standardization is applied only to the predictor variables, whereas it is also possible to apply it to the response variable.¹⁷

Table 2-16 Means and Standard Deviations for Fuel Properties, Based on Fuel Matrices for the Full and Reduced Designs

Reduced Desi							
Model Term		Full Design ¹					
		Mean	Standard deviation				
Ethanol (%)		10.31	7.88				
Aromatics (%)		25.63	10.02				
RVP (psi)		8.52	1.61				
T50 (°F)		190.61	28.58				
T90 (°F)		320.53	19.48				
etOH × etOH		0.96	0.80				
T50 × T50		0.96	0.74				
etOH × Arom		-0.037	0.98				
etOH × RVP		-0.10	1.00				
etOH × T50		-0.54	0.77				
etOH × T90		0.016	0.97				

Reduced Design ²						
Mean	Standard deviation					
11.02	8.06					
24.39	9.92					
197.00	23.45					
323.53	19.60					

Notes:

¹ Applies to models fit with data for 15 vehicles measured on 27 fuels.

² Applies to models fit with data for 5 or 15 vehicles measured on 11 fuels. Note that these models have no linear term for RVP and no 2nd order terms.

Table 2-17 Examples of One-Stage and Two-Stage Standardization for Three Test Fuels (1, 5 and 20)

Fuel	etOH	Arom	RVP	T50	T90	etOH	T50	etOH	etOH	etOH	etOH
	(vol.%)	(vol.%)	(psi)	(°F)	(°F)	×	×	×	×	×	×
	(1011/0)	(101170)	(101)	(1)	(-)	etOH	T50	Arom	RVP	T50	T90

Fuel Properties

1	10.03	15.4	10.07	148.9	300.2
5	0.00	34.7	6.95	237.0	300.0
20	20.31	15.2	6.70	162.7	298.7
Mean ¹	10.314	25.630	8.518	190.6	320.5
Std.					
Dev. ¹	7.880	10.015	1.611	28.6	19.5

One-Stage Standardized Values (Z) (Equation 9)

	$Z_{\rm e}$	$Z_{\rm a}$	$Z_{\rm r}$	Z_5	Z_9
1	-0.036	-1.021	0.963	-1.460	-1.044
5	-1.309	0.906	-0.973	1.623	-1.054
20	1.269	-1.041	-1.128	-0.977	-1.121

Mean ²			0.9630	0.9630	-0.0367	-0.0992	-0.5413	0.1633
Std.								
Dev. ²			0.8028	0.7398	0.9785	0.9996	0.7692	0.9728

Two-Stage Standardized Values (ZZ) (Equation 10, Equation 11)

	ZZ_{ee}	ZZ_{55}	ZZ_{ea}	ZZ_{er}	ZZ_{e5}	ZZ_{e9}
1	-1.198	1.578	0.075	0.065	0.772	0.022
5	0.935	2.260	-1.174	1.373	-2.058	1.401
20	0.805	-0.012	-1.313	-1.332	-0.907	-1.478

Notes:

Model Fitting

Throughout model fitting, the response variable was the natural logarithm transformation of the emissions results $(\ln Y)$, and the predictor variables were the one- or two-stage standardized fuel properties, as shown in Table 2-17. Thus, the model to be fit includes some subset of the 11 candidate terms shown in Equation 12.

$$\begin{array}{l} \ln Y = \beta_0 + \\ \beta_1 Z_e + \beta_2 Z_a + \beta_3 Z_r + \beta_4 Z_5 + \beta_5 Z_9 + \\ \beta_6 Z Z_{55} + \beta_7 Z Z_{ee} + \\ \beta_8 Z Z_{ea} + \beta_9 Z Z_{er} + \beta_{10} Z Z_{e5} + \beta_{11} Z Z_{e9} + \\ \varepsilon \end{array}$$
 Equation 12

A model containing all potential candidate terms is referred to as a "full model," whereas a model containing some subset of the candidate terms is referred to as a "reduced model." The goal of model fitting is to identify a reduced model by removing terms from the full model that do not contribute to fit.

¹ Mean and Standard Deviations of fuel properties for the entire fuel set. See Table 2-16.

² Mean and Standard Deviations of 2nd order terms values for the entire fuel set, constructed from the one-stage Z values.

Where the available data were sufficient, "mixed models" were fit, in which the terms listed in Table 2-16 were included as "fixed" terms. In addition, a "random intercept" was fit for each vehicle, which represents the high degree of variability contributed to the dataset by the vehicles measured. One way of understanding this distinction that the fuel properties are "fixed" because the fuels studied span the entire range of properties under study, and because the goal of the analysis is to estimate the effect of these parameters on the mean levels of emissions. On the other hand, "vehicle" is treated as a "random" factor because the sample of vehicles measured is but one of many samples that could have been measured. In the analysis, the emission levels of the specific vehicles are not of interest *per se*, but rather the degree of variability contributed to the analysis by the different vehicles. Analyses were performed using the MIXED procedure in the Statistical Analysis System (SAS®), version 9.2.¹⁸

When data were not sufficient for the mixed-model approach, models were fit by "Tobit regression." This technique was used when specific datasets were affected by low-end "censoring." For some measurements, the sample ostensibly obtained from the vehicle exhaust was lower than that attributable to background levels. In these cases, we assumed that a small but detectable mass was not measured accurately due to limitations in the sampling technique. In the Tobit model, the fitting method (maximum likelihood) is modified so as to compensate for the absence of the censored measurements. As with the mixed models, individual intercepts were fit for each vehicle; however, as the Tobit procedure does not distinguish "fixed" and "random" factors, vehicles were entered into the model as fixed factors (i.e., "dummy" variables). The Tobit models were fit using the LIFEREG procedure in SAS 9.2.¹⁹

Model fitting was conducted by backwards elimination, in which all terms in the full model were included at the outset. In fitting successive models, terms not contributing to fit were removed based on results of likelihood-ratio tests (LRT). Note that the LRT were used for model selection because all models were fit using "maximum-likelihood" (rather than "least-squares") methods.

Model fitting results for acetaldehyde, formaldehyde, acrolein and ethanol are shown in Table 2-18 through Table 2-21. Note that these four models represent "Bag 1" or "start" emissions on the LA92 cycle, based on datasets incorporating the full design. Also note that in fitting these models, an additional six terms beyond the original 11 design terms were included in the full models. These terms included one quadratic term (T90×T90), three interaction terms for aromatics, one interaction for RVP, and one interaction for the distillation parameters (T50×T90). However, none of these additional terms were retained as significant, with the single exception of the T50×T90 term.

In MOVES, emissions of toxics are estimated as fractions of volatile organic compounds in exhaust (VOC). To allow estimation of VOC, it was necessary to develop models for non-methane organic gases (NMOG). NMOG is equivalent to VOC, plus the mass of ethane and acetone. It is calculated in MOVES from non-methane hydrocarbons (NMHC) by correcting for the mass of oxygenated compounds not fully measured by the flame ionization detector used to determine NMHC. PA and CARB regulations set NMOG emission standards for motor vehicles, so NMOG is an important model output. The model representing start emissions for

-

^d Note that acetone was treated as negligible for purposes of these calculations.

NMOG, fit using the full design, is shown in Table 2-22. This model was fit using the same methods as that for total hydrocarbons (THC), as described in the Fuel Effects Report.²²

Table 2-18 Acetaldehyde (Bag 1): Coefficients and Tests of Effect for the Full and Reduced Models¹

1 401	L 2	-10 Acctan	ichyde (Dag	, 1). C	ocinciciit	s and 1 csts	, 0,	Effect for th	c Full allu	ixcuu	cu mo	ucis
Effect			Fuli	! Mode	el				Reduced	d Mode	el	
		Estimate	Std.Err.	d.f.	<i>t</i> -value	Pr>t		Estimate	Std.Err.	d.f.	t-	Pr>t
											value	
Intercept		-5.2324	0.08802	15	-59.4	0.000000		-5.2323	0.08785	15	-59.6	0.000000
Z_{e}		0.8250	0.01297	898	63.6	0.000000		0.8145	0.01020	898	79.9	0.000000
Z_a		0.03999	0.009279	898	4.31	0.000018		0.03484	0.008249	898	4.22	0.000027
Z_r		-0.03667	0.01297	898	-2.83	0.0048		-0.04170	0.008833	898	-4.72	0.000003
Z_5		0.09927	0.01826	898	5.44	0.000000		0.08670	0.01063	898	8.16	0.000000
Z_9		0.04235	0.01115	898	3.80	0.00016		0.03801	0.007764	898	4.90	0.000001
ZZ_{ee}		-0.1716	0.01548	898	-11.09	0.000000		-0.1669	0.007849	898	-21.3	0.000000
ZZ_{55}		0.07115	0.01314	898	5.42	0.000000		0.06665	0.007993	898	8.34	0.000000
ZZ_{ea}		0.03016	0.01304	898	2.31	0.021		0.01840	0.007777	898	2.37	0.018
ZZ_{er}		0.02020	0.008769	898	2.30	0.021		0.02194	0.007845	898	2.80	0.0053
ZZ_{e5}		-0.01614	0.01673	898	-0.965	0.33						
ZZ_{e9}		-0.01486	0.01072	898	-1.39	0.17						
						T	1		1		1	T
ZZ_{ar}		0.01738	0.01618	898	1.07	0.28						
ZZ_{a5}		0.004828	0.01729	898	0.28	0.78						
ZZ_{a9}		0.008759	0.008852	898	0.99	0.32						
ZZ_{99}		0.01270	0.01503	898	0.84	0.40						
ZZ_{59}		0.02718	0.01132	898	2.49	0.013		0.03959	0.008256	898	4.80	0.000002
ZZ_{r9}		-0.0206	0.009971	898	-2.07	0.039						
σ_{veh}^2		0.1154					-	0.1149				
σ_{ε}^2		0.08743						0.08850				

¹ See 9.2.2 and 8.7.3 in the Project Report. ¹⁹

Table 2-19 Formaldehyde (Bag 1): Coefficients and Tests of Effect for Full and Reduced Models¹

Effect		1) I Orma	_ •	Mode			Reduced Model					
		Estimate	Std.Err.	d.f.	<i>t</i> -value	Pr>t		Estimate	Std.Err.	d.f.	t-	Pr> <i>t</i>
											value	
Intercept		-5.9771	0.1498	15	-39.9	0.000000		-5.9771	0.1498	15	-39.9	0.000000
$Z_{ m e}$		0.2279	0.01234	898	18.5	0.000000		0.2299	0.009640	898	23.8	0.000000
Z_a		0.03528	0.008841	898	3.99	0.000071		0.02822	0.007979	898	3.54	0.00043
Z_r		-0.05202	0.01234	898	-4.21	0.000028		-0.04718	0.008457	898	-5.58	0.000000
Z_5		0.1577	0.01738	898	9.07	0.000000		0.1672	0.01001	898	16.7	0.000000
Z_9		0.1357	0.01064	898	12.7	0.000000		0.1302	0.007360	898	17.7	0.000000
ZZ_{ee}		-0.01498	0.01475	898	-1.02	0.31						
ZZ_{55}		0.05026	0.01251	898	4.02	0.000064		0.05262	0.008341	898	6.31	0.000000
ZZ_{ea}		0.02017	0.01241	898	1.63	0.10		0.01651	0.007340	898	2.25	0.025
ZZ_{er}		0.004100	0.008366	898	0.490	0.62						
ZZ_{e5}		-0.03686	0.01594	898	-2.31	0.021		-0.01627	0.008177	898	-1.99	0.047
ZZ_{e9}		0.02181	0.01023	898	2.13	0.033		0.02004	0.008838	898	2.27	0.024
	ı					1						
ZZ_{ar}		0.007384	0.01535	898	0.481	0.63						
ZZ_{a5}		-0.006739	0.01645	898	-0.41	0.68						
ZZ_{a9}		-0.01036	0.008437	898	-1.23	0.22						
ZZ_{99}		0.02104	0.01435	898	1.47	0.14						
ZZ_{59}		0.03974	0.01080	898	3.68	0.00025		0.03489	0.009322	898	3.74	0.00019
ZZ_{r9}		-0.003140	0.009498	898	-0.331	0.74						
σ_{veh}^2		0.3360					,	0.3358				<u> </u>
σ_{ε}^2		0.1395						0.1406				

¹ See 9.2.2 and Appendix L.3 in the Project Report. ¹⁶

Table 2-20 Acrolein (Bag 1): Coefficients and Tests of Effect for Full and Reduced Models¹

Effect			Mode	el			Reduced Model (FM8)				
	Estimate	Std.Err.	d.f.	t-	Pr>t		Estimate	Std.Err.	d.f.	t-	Pr>t
				value						value	
Intercept ²	-7.9337						-7.9338				
Z_{e}	0.2571	0.02638	15	9.74	0.000000		0.2476	0.02738	15	9.04	0.000000
Z_a	0.1149	0.02128	15	5.40	0.000074		0.1122	0.02184	15	5.14	0.00012
Z_r	-0.05815	0.01799	15	-3.23	0.0056		-0.0645	0.01364	15	-4.73	0.00027
Z_5	0.1979	0.03123	15	6.34	0.000013		0.1881	0.03554	15	5.29	0.000091
Z_9	0.2465	0.02979	15	8.28	0.000000		0.2488	0.03125	15	7.96	0.000000
ZZ_{ee}	-0.06009	0.01880	15	-3.20	0.0060		-0.08306	0.01392	15	-5.97	0.000026
ZZ_{55}	0.02735	0.01709	15	1.60	0.13						
ZZ_{ea}	0.01716	0.01838	15	0.93	0.37						
ZZ_{er}	0.01253	0.01404	15	0.89	0.39						
ZZ_{e5}	-0.09661	0.02096	15	-4.61	0.00034		-0.1185	0.02415	15	-4.91	0.00019
ZZ_{e9}	0.04178	0.01618	15	2.58	0.021		0.04618	0.01120	15	4.12	0.00091
						ı			l	l	1
ZZ_{ar}	0.02002	0.01562	15	1.28	0.22						
ZZ_{a5}	0.01127	0.01822	15	0.62	0.55						
ZZ_{a9}	-0.007484	0.01726	15	-0.43	0.67						
ZZ_{99}	0.0004162	0.01481	15	0.028	0.98						
ZZ_{59}	0.06274	0.01552	15	4.04	0.0011		0.05985	0.01271	15	4.71	0.00028
ZZ_{r9}	0.0002551	0.01709	15	0.015	0.99						
-2 1	0.2622						0.2620	ĺ			
σ_{veh}^{2}	0.3633						0.3629				
$\sigma_{arepsilon}^2$	0.03206						0.3213				

¹ See 9.2.2 and 8.7.4 in the Project Report¹⁶
² Not fit by the Tobit model, manually recalculated from intercepts for individual vehicles.

Table 2-21 Ethanol (Bag 1): Coefficients and Tests of Effect for Full and Reduced Models

1	ลม	ie 2-21 Euna	moi (dag	1): C	oemcien	ts and rest	5 U	I Ellect for ru	n and Nec	luceu	Models	•
Effect			Ful	l Mod	'el				Reduced	l Mod	el	
		Estimate	Std.Err.	d.f.	t-	Pr>t		Estimate	Std.Err.	d.f.	t-	Pr>t
					value						value	
Intercept ²				15				-4.9081				
Z_{e}		1.4759	0.07240	15	20.38	< 0.00001		1.4643	0.07115	15	20.56	< 0.00001
Z_a		-0.0067	0.04327	15	-0.16	0.88						
Z_r		-0.05004	0.04316	15	-1.16	0.26		-0.05990	0.02940	15	-2.06	0.057
Z_5		0.1050	0.03806	15	2.76	0.015		0.07188	0.02964	15	2.37	0.032
Z_9		-0.1261	0.03701	15	-3.47	0.0034		-0.09990	0.03574	15	-2.78	0.014
ZZ_{ee}		-0.4787	0.06014	15	-7.96	< 0.00001		-0.4967	0.05229	15	-9.51	< 0.00001
ZZ_{55}		0.1261	0.05018	15	2.51	0.024		0.1121	0.03826	15	2.90	0.011
ZZ_{ea}		-0.005952	0.03881	15	-0.15	0.88						
ZZ_{e5}		0.02820	0.05277	15	0.54	0.60						
ZZ_{e9}		0.0008509	0.06491	15	0.0090	0.99						
ZZ_{er}		0.03237	0.05103	15	0.64	0.53						
	- 1				ı						I	
ZZ_{a5}		0.03318	0.03212	15	1.04	0.32						
ZZ_{a9}		-0.01143	0.03461	15	-0.33	0.74						
ZZ_{99}		-0.5112	0.04523	15	-1.13	0.28						
ZZ_{59}		0.05311	0.04341	15	1.22	0.24						
ZZ_{ar}		0.04136	0.02855	15	1.45	0.17						
ZZ_{r9}		-0.008676	0.04644	15	-0.20	0.85						
$\sigma_{ m veh}^{2-1}$								0.1283				
$\sigma_{arepsilon}^2$		0.5697						0.05739				

 $^{^1\,\}mathrm{See}~9.2.2$ in the Project Report. 16 2 Not fit by the Tobit model, manually recalculated from intercepts for individual vehicles.

Table 2-22 NMOG (Bag 1): Coefficients and Tests of Effect for Full and Reduced Models¹

t-value

-10.48

6.027

9.424

-4.037

10.09

2.513

5.656

1.862

2.452

Pr > t

< 0.0001

< 0.0001

< 0.0001

< 0.0001

< 0.0001

0.012

< 0.0001

0.063

0.014

1 ab	ie <u>z-zz mwi</u>	UG (Dag .	1): C0	emcient	s and res	เร บ	I Ellect for	run and	Reduc	eu w
Effect		Ful	l Mod	el				Redu	ced Mo	odel
	Estimate	Std.Err.	d.f.	t-	Pr > t		Estimate	Std.Err.	d.f.	t-va
				value						
Intercept	-0.9520	0.09077	15	-10.49	< 0.0001		-0.9521	0.09089	15	-10
$Z_{ m e}$	0.07981	0.01326	941	6.02	< 0.0001		0.08019	0.01330	941	6.0
Z_a	0.08789	0.00929	941	9.46	< 0.0001		0.08782	0.00932	941	9.4
Z_r	-0.04595	0.01053	941	-4.36	< 0.0001		-0.04224	0.01046	941	-4.0
Z_5	0.1344	0.01329	941	10.12	< 0.0001		0.1345	0.01333	941	10
Z_9	0.01593	0.00925	941	1.72	0.0855					
ZZ_{ee}	0.04594	0.01760	941	2.61	0.00918		0.04432	0.01764	941	2.5
ZZ_{55}	0.07680	0.01336	941	5.75	< 0.0001		0.07579	0.01340	941	5.6
ZZ_{ea}	0.01635	0.00906	941	1.80	0.0714		0.01693	0.00909	941	1.8
ZZ_{er}	-	-	-	-	-					
ZZ_{e5}	0.04754	0.01893	941	2.51	0.0122		0.04653	0.01898	941	2.4
ZZ_{e9}	0.01961	0.00902	941	2.17	0.0300					
σ^2	0.1224						0.1224			

0.1224
0.07538

¹ See 9.1.2 in the Project Report¹⁶

Model Development under the Reduced Design

0.07538

As previously discussed, the "reduced design" involved the measurement of 11 fuels on 5 or 15 test vehicles, whereas the "full design" involved measurement of 27 fuels on 15 vehicles.

As shown in Table 2-15, measurements of two compounds in Bag 1, and all compounds in Bag 2, were performed under the reduced design. Supplementary analyses suggested that the reduced design was not adequate to support model fitting as described in Section 0 above. These results suggested that in these cases, full models retaining all four linear terms would perform as well or better than corresponding reduced models, many of which would retain only single terms. Thus, this sub-section presents results for full models under the reduced design.

Models representing start (Bag 1 on LA92) emissions are presented for benzene, 1,3-butadiene, non-methane organic gases (NMOG) and ethane in Table 2-23 through Table 2-26. These models were fit using subsets of data incorporating 15 vehicles measured over 11 fuels.

Similarly, models representing hot-running (Bag 2 on LA92) emissions are presented for acetaldehyde, formaldehyde, ethanol, NMOG and ethane in Table 2-27 through Table 2-31. These models were fit using subsets of data incorporating five vehicles measured over 11 fuels.

The development of these models is described in greater detail in sub-section 9.2.1 of the EPAct analysis report.¹⁶

Table 2-23 Benzene (Bag 1): Coefficients and Tests of Effect for the Full Model (Fit Under the Reduced Design, with 15 Vehicles, 11 Fuels)¹

(Fit Chaci	u	ic ixcuuccu i	ocsign, wi	1111	v chickes, i	i i ucis,						
Effect			Full Model									
		Estimate	Std.Err.	d.f.	t-value	Pr>t						
Intercept		-4.1019	0.1392	15	-29.48	< 0.0001						
$Z_{\rm e}$		-0.004685	0.03704	161	-0.126	0.90						
Z_a		0.4056	0.03389	161	11.97	< 0.0001						
Z_5		0.04142	0.03789	161	1.09	0.28						
Z_9		0.01133	0.03255	161	0.35	0.73						
$\sigma_{ m veh}^2$		0.2741										
σ_c^2		0.1873										

 $[\]frac{\partial \varepsilon}{\partial \varepsilon}$ See 9.2.2 and Appendix O.3 to the Project Report. ¹⁶

Table 2-24 1,3-Butadiene (Bag 1): Coefficients and Tests of Effect for the Full Model (Fit Under the Reduced Design, with 15 Vehicles, 11 Fuels)¹

The Chach the Reduced Design, with 13 venicles, 11 rucis,												
Effect			Full Model									
		Estimate	Std.Err.	d.f.	t-value	Pr>t						
Intercept		-5.8371	0.1235	15	-47.28	1.06×10 ⁻¹⁷						
$Z_{\rm e}$		-0.01729	0.03071	160	-0.56	0.57						
Z_a		0.02673	0.02730	160	0.98	0.33						
Z_5		0.01247	0.03031	160	4.11	0.000062						
Z_9		0.10036	0.02657	160	3.78	0.00022						
$\sigma_{ m veh}^2$		0.2192		•	•							
$\sigma_{arepsilon}^2$		0.1089										

¹See 9.2.2 in the Project Report. ¹⁶

Table 2-25 NMOG (Bag 1): Coefficients and Tests of Effect for the Full Models (Fit under the Reduced Design, 15 vehicles, 11 fuels)¹

(The under the recodered Besign, 18 venteres, 11 rueis)							
Effect	Full Model						
	Estimate	Std.Err.	d.f.	t-value	Pr> <i>t</i>		
Intercept	-0.8943	0.08668	15	-10.32	0.000000033		
Z_{e}	0.1040	0.01921	362	5.411	0.00000011		
Z_a	0.09435	0.01697	362	5.559	0.000000053		
Z_5	0.1527	0.01890	362	8.079	0.000000000		
Z_9	0.02127	0.01648	362	1.290	0.198		
$\sigma_{ m veh}^2$	0.1091			•			
$\sigma_{\rm s}^2$	0.08907						

¹See 9.2.2 in the Project Report. ¹⁶

Table 2-26 Ethane (Bag 1): Coefficients and Tests of Effect for the Full Models (Fit Under the Reduced Design, with 15 Vehicles, 11 Fuels)¹

(Fit Under the Reduced Design, with 15 vehicles, 11 Fuels)								
Effect		Full Model						
		Estimate	Std.Err.	d.f.	t-value	Pr>t		
Intercept		-4.308	0.09833	15.0	-43.81	2.84×10 ⁻¹⁷		
Z_{e}		0.1204	0.02075	160	5.805	3.37×10 ⁻⁸		
Z_a		-0.1728	0.01844	160	-9.373	6.51×10 ⁻¹⁷		
Z_5		0.2169	0.02047	160	10.59	3.30×10 ⁻²⁰		
Z_9		0.09531	0.01795	160	5.311	3.60×10 ⁻⁷		
$\sigma_{ m veh}^2$		0.1407						
$\sigma_{\mathfrak{s}}^2$		0.04970						

¹ See 9.2.2 in the Project Report. ¹⁶

Table 2-27 Acetaldehyde (Bag 2): Coefficients and Tests of Effect for the Full Models
(Fit Under the Reduced Design, with 5 Vehicles, 11 Fuels)¹

The United the Reduced Design, with 5 vehicles, 11 Fuels)							
Effect		Full Model					
		Estimate	Std.Err.	d.f.	<i>t</i> -value	Pr> <i>t</i>	
Intercept		-9.4189	0.1177	5	-80.1	0.000000	
Z_{e}		0.1520	0.06080	58	2.50	0.0152	
Z_a		0.07991	0.05279	58	1.51	0.136	
Z_5		-0.02997	0.05957	58	-0.503	0.617	
\mathbb{Z}_9		-0.07836	0.05153	58	-1.52	0.134	
$\sigma_{ m veh}^2$		0.05654					
$\sigma_{arepsilon}^2$		0.3814					

¹ See 9.2.2 and Appendix K.3 to the Project Report. ¹⁶

Table 2-28 Formaldehyde (Bag 2): Coefficients and Tests of Effect for the Full Model (Fit Under the Reduced Design, with 5 Vehicles, 11 Fuels)¹

		0 /				
Effect	Full Model					
	Estimate	Std.Err.	d.f.	<i>t</i> -value	Pr>t	
Intercept	-8.6574	0.1372	5.01	-63.10	< 0.00001	
$Z_{\rm e}$	0.08456	0.05937	58.04	1.424	0.16	
Z_a	0.01575	0.05154	58.05	0.306	0.76	
Z_5	0.01863	0.05815	58.03	0.320	0.75	
Z_9	-0.08138	0.05031	58.16	-1.62	0.11	
$\sigma_{ m veh}^2$	0.08205			•		
$\sigma_{\rm s}^2$	0.3762					

See 9.2.2 and Appendix L.4 to the Project Report. 16

Table 2-29 Ethanol (Bag 2): Coefficients and Tests of Effect for the Full Model (Fit Under the Reduced Design, with 5 Vehicles, 11 Fuels)¹

Fit Under the Reduced Design, with 5 vehicles, 11 Fuels)						
Effect		Full Model				
		Estimate	Std.Err.	d.f.	<i>t</i> -value	Pr>t
Intercept ¹		-9.3072	0.6333	5	-15.45	0.000021
Z_{e}		0.9233	0.2824	5	3.27	0.022
Z_a		-0.3772	0.28499	5	-1.32	0.24
Z_5		01910	0.2091	5	-0.091	0.93
Z_9		-0.3017	0.2416	5	-1.25	0.27
$\sigma_{\mathrm{veh}}^{2}$		0.3707		•		•
σ_{ε}^2		1.0889				

¹ See 9.2.2 and Appendix N.4 to the Project Report. ¹⁶

Table 2-30 NMOG (Bag 2): Coefficients and Tests of Effect for the Full Model
(Fit Under the Reduced Design with 5 Vehicles, 11 Fuels)¹

(Fit Under the Reduced Design, with 5 venicles, 11 Fuels)						
Effect		Full Model				
		Estimate	Std.Err.	d.f.	<i>t</i> -value	Pr> <i>t</i>
Intercept ¹		-4.777	0.4784	5	-9.99	0.00017
Z_{e}		0.01778	0.03574	124	0.497	0.62
Z_a		0.03320	0.03117	124	1.07	0.29
Z_5		0.04258	0.03494	124	1.22	0.23
Z_9		0.09051	0.03038	124	2.98	0.0035
$\sigma_{ m veh}^{2}$		1.1405				
$\sigma_{arepsilon}^2$		0.1026				
	- 1	~ ~ ~ ~ .			. 16	

¹ See 9.2.2 in the Project Report. ¹⁶

Table 2-31 Ethane (Bag 2): Coefficients and Tests of Effect for the Full Model (Fit Under the Reduced Design, with 5 Vehicles, 11 Fuels)¹

(Fit Under the Reduced Design, with 5 vehicles, 11 Fuels)						
Effect		Full Model				
		Estimate	Std.Err.	d.f.	<i>t</i> -value	Pr>t
Intercept ¹		-7.724	0.7325	5	-10.54	0.00013
$Z_{ m e}$		0.07345	0.05873	57	1.251	0.22
Z_a		-0.1260	0.05151	57	-2.447	0.018
Z_5		0.1815	0.05727	57	3.168	0.0025
Z_9		0.1322	0.04994	57	2.647	0.010
$\sigma_{ m veh}^{2}$		2.6712				_
$\sigma_{arepsilon}^2$		0.1476				

¹ See 9.2.2 and Appendix Q.4 to the Project Report. ¹⁶

2.1.1.2.2 Application of EPAct Statistical Models in MOVES

We estimate the emissions of VOC toxic compounds as a fraction of emissions for VOC, on the same fuel. To model the behavior of the fraction with respect to changes in fuel properties, it was necessary to develop models for NMOG and ethane, as well as the toxics, because VOC is estimated as NMOG minus ethane.^e

^e In MOVES, VOC is typically calculated as NMOG – ethane – acetone, but for this purpose, acetone was considered negligible, and was not subtracted.

The models generated using EPAct results allow estimation of emissions effects related to the five fuel properties included in the study design: ethanol content (vol. percent), aromatics content (vol percent), RVP (psi), T50 (°F) and T90 (°F), as well as selected interaction terms among these five parameters.

The statistical models generated from the EPAct data follow the general structure shown in Equation 13 below, which uses the model for acetaldehyde as an example (see Table 2-18). Note that the subsets of the potential terms vary by emission and process, depending on the results of model fitting, as described in the previous two sub-sections.

Emissions (g/mi) =
$$e^{X\beta}$$

= $exp\begin{pmatrix} \beta_0 + \beta_e Z_e + \beta_a Z_a + \beta_r Z_r + \beta_5 Z_5 + \beta_9 Z_9 + \\ \beta_{ee} Z Z_{ee} + \beta_{55} Z Z_{55} + \\ \beta_{ea} Z Z_{ea} + \beta_{er} Z Z_{er} + 0.5 (s_{veh}^2 + s_{\varepsilon}^2) \end{pmatrix}$ Equation 13
= $exp\begin{pmatrix} -5.23 + 0.814 Z_e + 0.0348 Z_a - 0.0417 Z_r + 0.0867 Z_5 + 0.0380 Z_9 - \\ 0.1669 Z Z_{ee} + 0.0667 Z Z_{55} + \\ 0.0184 Z Z_{ea} + 0.0219 Z Z_{er} + 0.5 (0.1149 + 0.08850) \end{pmatrix}$

Where the data were sufficient, two sets of exhaust fuel effect coefficients were employed for each pollutant; one set representing cold start emissions and a second set representing hot-running emissions. In some cases, fuel effects estimated for these two processes differed substantially, as the effects of fuel properties on start emissions are dominated by changes in combustion and catalyst warm-up, while the impact of gasoline properties on running emissions is dictated by catalyst efficiency when fully operational. Thus, using convenient matrix notation, the expressions $\mathbf{X}\boldsymbol{\beta}_{toxic}$, $\mathbf{X}\boldsymbol{\alpha}_{NMOG}$ and $\mathbf{X}\boldsymbol{\theta}_{ethane}$ represent models for a selected toxic compound, NMOG and ethane, respectively, calculated by applying Equation 13 to each compound for a specified fuel. The toxic emissions as a fraction of VOC emissions (f_{toxic}) are given by

Toxic Fraction =
$$f_{\text{toxic}} = \frac{e^{X\beta_{\text{toxic}}}}{e^{X\alpha_{\text{NMOG}}} - e^{X\theta_{\text{ethane}}}}$$
 Equation 14

For all compounds, the calculation shown in Equation 14 is recorded in the MOVES GeneralFuelRatioExpression table. In calculating toxic fractions, we elected to use models for NMOG and ethane fit using study designs and datasets similar to those for the toxic compounds. In other words, if the toxic model was fit with the reduced design, we combined it with the NMOG and ethane models also fit with the reduced design. We followed this approach to prevent the calculation and propagation of artifacts in the estimated fractions resulting from differing levels of information and complexity in the numerator and denominator in Equation 14. In this context, we considered it important to apply "information parity" to the toxic model in the numerator and the NMOG model in the denominator, as the vast majority of VOC mass is represented by NMOG, with ethane comprising only a small fraction.

Note that for three compounds in Bag 2, levels of "left censoring," were high enough that modeling was not considered feasible. Again, "censoring" occurs when background levels of the compounds under study were as high as or higher than levels ostensibly measurable in vehicle

exhaust. Estimation of "simple" toxic fractions for these compounds is covered in the following sub-section.

Estimating Simple Fractions of VOC for Running Emissions

As noted in Table 2-15, the bag 2 data were not sufficient to fit models for running emissions for three compounds: acrolein, benzene and 1,3-butadiene. The bag 1 data for these pollutants was sufficient to estimate start emission adjustments as discussed in the previous subsections. Therefore, running emissions were represented as "simple" (constant) fractions of VOC, with values derived from the available data, as shown in Table 2-32.

Table 2-32 Simple Fractions of VOC for Running Emissions for 2001 and Later Model Year Gasoline Vehicles

Compound	Simple Fraction
Acrolein	0.00077
Benzene*	0.047
1,3-butadiene	0.0

^{*}Benzene fractions are further adjusted according to the benzene content of the fuel as discussed in Section 2.1.1.2.3

These values were derived as "ratios of means" (ROM), in which the toxic and VOC values were averaged first by vehicle and then across vehicles, as described below. The ROM approach is generally preferred as it provides an unbiased estimator of the true fraction as the sample size increases.²³

For benzene, results were available for four vehicles, differing widely in their benzene and VOC levels, and also in numbers of available measurements, as shown in Table 2-33. The averaging was performed in two steps so that the vehicle(s) with the greatest numbers of measurements would not dominate the overall mean. In the first step, the benzene and VOC values were averaged for each vehicle. In the second step, the four vehicle means were averaged to give an overall mean. Finally, the overall mean for benzene was divided by that for VOC to give a simple ratio estimator for benzene as a fraction of VOC.

Table 2-33 Benzene (Running): Derivation of a Ratio-of-Means Estimator for Benzene as a Fraction of VOC1

Vehicle	n	Benzene (mg)	VOC (mg)	Ratio of means (ROM) ²
Corolla	2	0.054	2.269	
F150	10	2.224	28.427	
Impala	3	0.108	10.670	
Silverado	4	0.294	16.216	
All vehicles	4	0.670	14.396	0.0465

¹ Benzene fractions are adjusted according to the benzene content of the fuel as discussed in Section 2.1.1.2.3

The VOC fraction for acrolein was derived similarly (Table 2-34). For this compound results were available for five vehicles. Values for acrolein are considerably lower than for benzene, so results are expressed in µg, rather than mg. The resulting fraction is two orders of magnitude lower than that for benzene.

² This value is a simple average of the means for all four vehicles, as listed above.

Vehicle	n	Acrolein (μg)	VOC (µg)	Ratio of means (ROM)
Civic	3	5.42	3,038.9	
Corolla	5	2.89	2,929.6	
F150	5	8.36	24,321	
Impala	6	8.02	10,408	
Silverado	10	19.66	17,192	
All vehicles	5	8.87	11,578	0.00077

¹ This value is a simple average of the means for all five vehicles, as listed above.

For 1,3-butadiene in hot-running operation, measurements were extremely low; in fact, we considered the dataset so heavily affected by "left-censoring" that we did not consider it adequate for either model fitting or development of ratio estimators. Accordingly, for modeling purposes, we have adopted an assumption that this compound is not emitted during hot-running operation, i.e., the ROM estimator is 0.0.

2.1.1.2.3 Additional Fuel Adjustments for Benzene and 1,3-Butadiene

For two compounds, benzene and 1,3-butadiene, additional refinements were applied to supplement the study design of the EPAct fuel set. These adjustments are applied to both start and running emissions.

For benzene, the issue is that the fuel matrix in the EPAct test program included aromatics generally, but not benzene specifically. As we considered it inadequate to model benzene in exhaust without explicitly accounting for benzene levels in fuel, we developed a "post-EPAct model" refinement using data external to the EPAct program. In this case, the source was a program conducted in support of the 2007 MSAT2 rule. This program performed measurements on nine Tier-2 certified vehicles on fuels with benzene levels ranging from 0.6 to 1.1 percent by weight. With benzene represented as a fraction of VOC (as in Equation 14) denoted as fbenzene, a value modified to account for benzene levels in different fuels ($f^*_{benzene}$) is calculated as shown in Equation 15 where $x_{benzene}$ is the benzene level for the fuel modeled (weight percent), A is the mean benzene level in the EPAct exhaust program fuel set (0.66 weight percent), and B is an empirical coefficient, taking a value of 0.24.

$$f_{\text{benzene}}^* = [(x_{\text{benzene}} - A) \cdot B \cdot f_{\text{benzene}}] + f_{\text{benzene}}$$
 Equation 15

Similarly, given the importance of olefins to estimation of emissions for 1,3-butadiene, and that the EPAct exhaust program study design did not incorporate olefins as a factor, we considered it appropriate to develop a post-EPAct model adjustment explicitly accounting for olefin level. This adjustment was derived by varying olefin levels in the Complex Model and fitting a polynomial trend to the results. Starting with an unadjusted toxic fraction for 1,3-butadiene (f_{buta}), the modified fraction f_{buta}^* is calculated using Equation 16, in which x_{olefin} is the olefin level, and A, B, C and D are coefficients, taking values of 0.000008, 0.0002, 0.0069 and 0.008823, respectively.

$$f_{\text{buta}}^* = f_{\text{buta}} \left(\frac{Ax_{\text{olefin}}^2 + Bx_{\text{olefin}} + C}{D} \right)$$
 Equation 16

Equation 16 is applied to the estimated 1,3-butadiene fraction for start emissions estimated from the EPAct equations in Table 2-24. MOVES does not apply similar fuel adjustments to the 1,3-butadiene running exhaust emissions because they are estimated to be zero (Table 2-32).

2.1.1.2.4 Additional Air Toxics Estimated from EPAct Speciation Profiles

As summarized in Table 2-11, and analogous to what was done for 2000-and-earlier vehicles, we used a simpler approach for the seven hazardous air toxics listed in Table 2-35.

For fuel blends with 0 percent, 10 percent and 15 percent ethanol, composite speciation profiles developed from the results of EPAct (Phase 1) were used to develop toxic fractions of VOC for the hazardous air toxics listed in Table 2-35. These profiles were based on averaging results of tests from 3 vehicles. Toxic fractions for E10 are used for all gasolines containing ethanol levels of 5 vol. percent or greater. For fuel blends containing 20 percent ethanol, fractions were developed using a composite speciation profile from the EPAct (Phase 3) program. The fractions are also presented in Table 2-35. The values shown in Table 2-35 are stored in the database table minorHAPRatio (see Table 2-10).

Table 2-35 Toxic Fractions of VOC for Selected Compounds, Representing Model Years 2001 and Later

1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				
Pollutant (pollutantID) ¹				
2,2,4-Trimethylpentane (40)				
Ethyl Benzene (41)				
Hexane (42)				
Propionaldehyde (43)				
Styrene (44)				
Toluene (45)				
Xvlene(s) (46)				

Fuel Blends (Gasoline and Ethanol)					
0% (E0)	10% (E10) ²	15% (E15)	20% (E20) ³		
0.03188	0.01227	0.02198	0.004625		
0.01683	0.01660	0.01568	0.022199		
0.002790	0.02911	0.0110	0.02497		
0.00122	0.00054	0.0005984	0.0006607		
0.00085	0.00083	0.004588	0.004096		
0.07542	0.07440	0.0727	0.09646		
0.06127	0.06047	0.06902	0.09302		

¹ For fuels containing 0-20 percent ethanol, fractions for ethanol, benzene, acetaldehyde, formaldehyde, 1,3-butadiene, and acrolein were estimated using methods described in the previous subsections.

² Values also applied for fuels containing 5 percent and 8 percent ethanol, (E5 and E8)

³MOVES versions after MOVES2014a do not estimate emissions for vehicles using E20 fuel, but these fractions currently remain in the MOVES database.

^f Phase 1 testing was done using fuels more representative of in-use fuels, in contrast to the orthogonal matrix used for EPAct Phase 3.

2.1.2 Vehicles Operating on Fuel Blends Containing 70-100 Percent Ethanol

Flexible-fueled vehicles (FFVs) may operate on gasoline, or on "E85" fuel, which is a mix of gasoline with high levels of ethanol (typically 70-85%). Within this sub-section, we further delineate the methods and data used for estimating hazardous air pollutants in vehicles from model years 2000 and earlier (Section 2.1.2.1), 2001 and later model year vehicles (Section 2.1.1.2), and data that applies to all model year vehicles (Section 2.1.2.3). In this section, we distinguish "major HAPs" (benzene; ethanol; 1,3-butadiene, formaldehyde, acetaldehyde and acrolein) which have toxic ratios that can be a function of fuel properties. Other toxic VOCs ("minor HAPs": 2,2,4-trimethylpentane; ethyl benzene; hexane; propionaldehyde; styrene, toluene, and xylene(s)) use toxic ratios that vary by fuel subtype, but not as a function of fuel properties.

2.1.2.1 2000 and Earlier Model Year Vehicles

For major HAP emissions for vehicles in model years 2000 and earlier operating on fuel blends containing 70-100 percent ethanol are estimated as fractions of VOC. There are few pre-2000 FFVs, and little data on their emissions using E85 fuel. Thus, the toxic fractions were derived from data from four newer flexible-fuel vehicles running on E85 gasoline, collected during the EPAct program (Phase 3) as displayed in Table 2-36. Since no measurements were obtained on an E70 blend (more typically used in winter) or blends above E85, the same toxic-to-VOC fractions are used for all ethanol-gasoline blends containing 70-100 percent ethanol. These ratios are applied to older technology (2000 and earlier vehicles), even though data were collected from Tier 2 vehicles. The 2000 and earlier HAP emission rates are stored in the database table "ATRatioNonGas" (see Table 2-37).

Table 2-36 E70/E85 Major HAP VOC Fraction for 2000 and Earlier Model Year Vehicles

Pollutant (pollutantID)	Toxic Fraction
Benzene (20)	0.0170
Ethanol (21)	0.3724
1,3-butadiene (24)	0.0011
Formaldehyde (25)	0.0291
Acetaldehyde (26)	0.1644
Acrolein (27)	0.0010

Table 2-37 Description of the Database Table "ATRatioNonGas," as Applied to Light-Duty Vehicles

Field	Description	RelevantValues
Ficiu	Description	Refevant values
polProcessID	Identifies the pollutant (1st two digits and Emissions Process (last two digits).	Pollutants are identified in the table above; Relevant processes include: "Running Exhaust" (processID = 1) "Start Exhaust" (processID = 2) "Extended Idle Exhaust" (processID = 90) "Auxiliary Power Exhaust" (processID = 91)
sourceTypeID	Identifies types of vehicles, classified by function	Motorcycle (11) Passenger Car (21) Passenger Truck (31) Light Commercial Truck (32)
fuelSubTypeID	Identifies specific fuel classes within the fuelTypeID	51 = "Ethanol (E85)" 52 = "Ethanol (E70)"
modelYearGroupID	Identifies a set of model years covered by a specific value of atRatio.	In this table, the first and last model year of the group are concatenated, such as "19601970", or "20302030".
atRatio	Fraction, or "ratio" of the toxic relative to total VOC.	
atRatioCV	"Coefficient of Variation of the Mean" or "relative standard error" of the atRatio.	
dataSourceID	Indicates source data and methods used to estimate atRAtio.	

2.1.2.2 2001 and Later Model Year Vehicles

For major HAPs in 2001 and later model year vehicles using E-85 fuels, we conducted a more comprehensive analysis than for the older model year vehicles. Instead of deriving toxic fractions of VOC, we developed adjustment factors that were compatible with the EPAct toxic ratios derived for gasoline 2001 and later model year vehicles discussed in the Section 2.1.1.2. The toxic adjustment factors were developed based on the analysis of EPAct (Phase 3) program, National Renewable Energy Laboratory (NREL) E40²⁹, Coordinating Research Council (CRC) E-80³⁰, and the PM Speciation Program. All programs measured emissions from LA92 test cycle on both E10 and E85, except CRC E-80 which tested E6 and E85. Only the vehicles tested on both E10 (E6) and E85 were included in the analysis. Numbers of vehicles in each program are summarized in Table 2-38.

Table 2-38 Numbers of Vehicles Included in the Analysis of Major HAPs

Test Program	Number of Vehicles
EPAct (phase 3)	4
NREL E40	9
CRC E-80	7
PM Speciation	2

Consistent emission trends were observed across datasets; thus, the datasets were pooled to examine the effect of E85 on emissions compared to E10. First, the test of significance of differences between E10 and E85 was performed using Student's paired *t*-tests. Next, when statistically significant differences in emissions between E10 and E85 were evident, the adjustment factors were calculated using Equation 17. The adjustment factor was set to one when the differences in emissions were not statistically different (i.e., acrolein).

$$E85 \ adjustment \ factor = \frac{\frac{Toxics_{E85}}{VOC_{E85}}}{\frac{Toxics_{E10}}{VOC_{F10}}}$$
 Equation 17

The resulting adjustment factors are shown in Table 2-39. As expected, ethanol and the aldehydes (formaldehyde, acetaldehyde) have higher toxic ratios than the E10-fueled vehicles.

Table 2-39 E70/E85 Adjustment Factors for Major HAPs for 2001 and Later Model Year Vehicles

Pollutant (pollutantID)	Adjustment Factor for E70/E85
Benzene (20)	0.6672
Ethanol (21)	7.587
1,3-butadiene (24)	0.2167
Formaldehyde (25)	1.572
Acetaldehyde (26)	7.126
Acrolein (27)	1

The toxic fractions for the pollutants in Table 2-39 are calculated using Equation 18.

E85 Toxic Fraction =
$$f'_{toxic} \times E85$$
 adjustment factor
$$= \left(\frac{e^{X'\beta_{toxic}}}{e^{X'\alpha_{\text{NMOG}}} - e^{X'\theta_{\text{ethane}}}}\right) \times E85 \text{ adjustment factor}$$
Equation 18

The f'_{toxic} uses the same equations and coefficients as were derived for MY 2001+ gasoline vehicles discussed in Section 2.1.1.2 (including the adjustments for benzene and 1,3-butadiene). The difference between f'_{toxic} used for E85 vehicles and the f_{toxic} used for gasoline vehicles is that f'_{toxic} uses a combination of fuel properties from E85 and E10 fuels as shown in Table 2-40. As explained in the MOVES Fuel Effects Report²², MOVES uses the E10 fuel properties for ethanol content, aromatics, T50 and T90 from E10 fuels (stored in the E10FuelProperties table), because these fuel effects are obtained from the EPAct program, and the fuel equations are not intended to be used for the E85-specific levels of ethanol content, aromatics, T50 and T90, which are outside the bounds of the fuel properties included in the sample design. In addition, the E85 adjustment factors are intended to account for the differences between E85 and E10 emissions,

not the fuel effects from the EPAct study. Olefin content is also used from the E10 fuel properties table, because the olefin content from E85 fuel is significantly lower than typical gasoline fuels, and the 1,3-butadiene adjustments were developed from the Complex Model on gasoline fuels, not E85 fuels (Section 2.1.1.2.3). MOVES accounts for the differences in 1,3-butadiene for E85 fuels using the E85 adjustment factor (Table 2-39).

For RVP, benzene and sulfur levels, we use the E85-specific values because these fuel properties are within the range of gasoline fuels (RVP) or were not accounted for in the EPAct fuel design (benzene and sulfur), and we believe it is appropriate to apply the E85 specific-fuel properties.

Table 2-40 Source of Fuel Properties (E85 or E10) used to Estimate Toxic Fractions for MY 2001 and Later E85 Vehicles

Ethanol Volume	RVP	Sulfur Level	Benzene Content	Aromatic Content	Olefin Content	T50	Т90
E10 (10%)	E85	E85	E85	E10	E10	E10	E10

Equation 18 and the accompanying E85 adjustment factors are stored in the MOVES *GeneralFuelRatioExpression* table to estimate emissions for the six air toxics pollutants that are a function of fuel properties ("major HAPs"). Because the E85 toxic fractions are calculated using the fuel properties of RVP, sulfur and benzene content from E85 fuels, the resulting ratios between toxic emission rates from E85 and E10 fueled vehicles in MOVES are slightly different than the adjustment factors shown in Table 2-39. In addition, the VOC emissions from E85-fueled and E10-fueled vehicles (to which the toxic fractions apply) are slightly different due to the lower sulfur level of E85 fuel as discussed in the MOVES3 Fuel Effects Report.²²

2.1.2.3 Air Toxics Fractions that Apply to All Model Year Vehicles

Fractions for the remaining "minor HAPs" air toxic compounds modeled in MOVES were developed from the four flexible-fuel vehicles tested during the EPAct program (Phase 3) running on fuels containing 70-100 percent ethanol. The E85 toxics are consistent with the speciation profile developed from the same test data.²¹. As stated earlier, the vehicles were tested on a single E85 gasoline fuel. These ratios are applied to older technology (2000 and earlier vehicles) as well as the modern technology vehicles in the test program; thus, while there are few pre-2000 FFVs, there is more uncertainty in emission estimates for older technology vehicles running on high ethanol blends than for newer vehicles. As expected, the toxic fractions for the E85 vehicles shown in Table 2-41 are lower than E10-fueled vehicles for both pre-2001 MY (Table 2-9) and MY 2001 and later vehicles (Table 2-35), with the exception of propionaldehyde. The toxic fractions for the pollutants shown in Table 2-41 are stored in the minorHAPRatio table (see Table 2-10).

Table 2-41 Toxic Fractions of VOC for Vehicles Running on E70/E85 for All Model Year Vehicles

Pollutant (pollutantID)	Toxic Fraction of VOC
2,2,4-Trimethylpentane (40)	0.0078
Ethyl Benzene (41)	0.0055
Hexane (42)	0.0045
Propionaldehyde (43)	0.0025
Styrene (44)	0.0003
Toluene (45)	0.0177
Xylene(s) (46)	0.0185

2.2 Polycyclic Aromatic Hydrocarbons (PAHs)

Emissions of PAHs are estimated using fractions similar to that used for VOCs as described in the previous section. However, for PAHs, the process is complicated by the fact that exhaust and crankcase emissions of these compounds are emitted in both the gaseous and particulate phases. Accordingly, emissions in the gaseous phase are estimated as fractions of total VOC, and emissions in the particulate phase as fractions of organic carbon $\leq 2.5 \, \mu m$ (OC_{2.5}). We discuss the derivation of PAH fractions for vehicles operating on gasoline containing low ethanol percentages (Section 2.2.1) and high-ethanol percentages (Section 2.2.2).

2.2.1 Vehicles Operating on Fuel Blends Containing 0-15 Percent Ethanol

The PAH emission fractions for gasoline vehicles are estimated from a set of 99 vehicles selected for chemical speciation that were measured in the Kansas City Light-duty Vehicle Emissions Study (KCVES).³² Each vehicle was measured on the LA92 driving cycle that includes both start and running operation. For each vehicle, emissions of THC and particulate matter 2.5 microns in diameter or less (PM_{2.5}) were measured. Fleet-average fractions of PAH/THC and PAH/PM_{2.5} were calculated with each sample weighted by total emissions,^g vehicle-miles traveled (VMT), and an equal weight between summer and winter. We used a VOC/THC fraction of 0.86 developed from the total organic-gas speciation profile developed from the Kansas City program (8750a) to estimate PAH/VOC fractions. Because the PAH measurements from the LA92 cycle include both start and running emissions, we use the same VOC fractions for both start and running PAH emissions. We adjusted the PAH/PM_{2.5} fraction by the fraction of OC measured in the start (42.6 percent) and running emission processes (55.7 percent) to produce PAH/OC_{2.5} emission fractions. Because OC/PM fractions differ for start and running, we have separate PAH/OC toxic fractions for start and running.

The partitioning of PAH emissions between gaseous and particulate phases is assigned on the basis of average temperature and dilution conditions at the time of measurement, i.e., in the sample train and constant-volume sampler. Thus, the partitioning reflected in the emission fractions does not reflect cooling and dilution occurring in the "real world" after the exhaust leaves the tailpipe. The sampling conditions set forth in EPA regulations for particulate and

_

g Each sample contained emissions from one to five vehicles.

hydrocarbon measurement differ for light-duty and heavy-duty vehicles, which affects the phase partitioning of PAH emissions obtained from both engine types. In preparing inputs for MOVES, we developed one set of phase allocation factors for gasoline sources and another for diesel sources in order to streamline data processing, and to be consistent with the measurement conditions reflected in the PAH measurements.

The allocations of PAHs into gaseous and particulate phases for gasoline vehicles was based on measurement samples analyzed by Desert Research Institute (DRI) on a subset of vehicles in the KCVES that were measured with dilution air at both low and high dilution temperatures.³³ One of the purposes of this follow-up study was to examine the impact of sampling conditions on PAH emission measurements. DRI measured PAH species with Teflon-impregnated glass filters (TIGF) and backup glass cartridges with Amberlite XAD-4 adsorbent resins over the LA92 cycle. Relative concentrations of individual PAH were measured on the TIGF and the XAD with sampling line and dilution temperatures of 20°C and 47°C for four composite samples, with each composite sample containing one to three vehicles. Table 2-42 reports the TIGF/XAD phase allocation factors measured at 47°C (which was the measurement temperature for the Kansas City Light-duty Vehicle Emissions Study), for the composite sample referred to as the 'mediumemitters.' This class contained a 1989 Camry and 1992 Voyager. In MOVES, we used the PAH phase-partitioning of this sample to estimate the relative gas and particle portioning of all gasoline-source emissions. Clearly, this sample may not adequately represent phase-partitioning of PAH emissions from the current in-use fleet; however, it was deemed the most representative of the breadth of gasoline vehicles sampled in the KCVES. Note that the PAH species partitioning was heavily dependent on molar mass (molecular weight); compounds with lighter molar masses (e.g., naphthalene) were measured almost entirely in the gaseous phase, whereas compounds with heavier molar masses were measured almost entirely in the particulate phase (e.g., dibenzo(a,h)anthracene).

Table 2-42 Gasoline PAH Phase Allocation Factors

PAH species	Molar Mass (g/mol)	Phase Fraction	
	(8')	Gaseous	Particulate
Naphthalene	128	0.9996	0.0004
Acenaphthylene	152	0.9985	0.0015
Acenapthene	154	1.0000	0.0000
Fluorene	166	1.0000	0.0000
Anthracene	178	0.9915	0.0085
Phenanthrene	178	0.9953	0.0047
Fluoranthene	202	0.9822	0.0178
Pyrene	202	0.9831	0.0169
Benz(a)anthracene	228	0.6721	0.3279
Chrysene	228	0.7307	0.2693
Benzo(a)pyrene	252	0.0426	0.9574
Benzo(b)fluoranthene	252	0.5546	0.4454
Benzo(k)fluoranthene	252	0.5546	0.4454
Benzo(g,h,i)perylene	276	0.0000	1.0000
Indeno(1,2,3-cd)pyrene	276	0.0000	1.0000
Dibenzo(a,h)anthracene	278	0.0000	1.0000

The PAH/VOC and PAH/OC emission fractions used in MOVES are calculated by multiplying the PAH/VOC, and PAH/OC fractions calculated from KCVES by the gas/particle partitioning factors in Table 2-42. The calculation is displayed with Equation 19 and Equation 20 for each PAH, i= 1:16.

$$\frac{PAH_{i}}{VOC} \text{(Table 2-43)} = \frac{PAH_{i}}{VOC} (KCVES) \times Gaseous \, Fraction_{i} \, \text{(Table 2-42)}$$

$$\frac{PAH_{i}}{OC} \text{(Table 2-43)}$$

$$= \frac{PAH_{i}}{OC} (KCVES) \times Particulate \, Fraction_{i} \, \text{(Table 2-42)}$$

Within MOVES, the PAH fractions in Table 2-43 are applied to all gasoline fuels with ethanol content less than 20 percent. In the MOVES database, these fractions are stored in two tables. Fractions for the gaseous and particulate phases are stored in the tables pahGasRatio and pahParticleRatio, respectively. The two tables have the same structure, which is presented in Table 2-43.

Table 2-43 Toxic Fractions for PAH Compounds, in Gaseous and Particulate Phases for Gasoline Vehicles Fueled with Ethanol Content < 20 percent

Species	Gaseous Phase (PAH/VOC)	Particulate Phase (PAH/OC2.5)	
		Start	Running
Naphthalene	2.07×10 ⁻³	1.68×10 ⁻⁴	1.29×10 ⁻⁴
Acenaphthylene	1.81×10 ⁻⁴	5.01×10 ⁻⁵	3.83×10 ⁻⁵
Acenaphthene	3.99×10 ⁻⁵	0.0	0.0
Fluorene	8.08×10 ⁻⁵	0.0	0.0
Anthracene	3.35×10 ⁻⁵	5.19×10 ⁻⁵	3.97×10 ⁻⁵
Phenanthrene	2.14×10 ⁻⁴	1.81×10^{-4}	1.39×10 ⁻⁴
Fluoranthene	5.60×10 ⁻⁵	1.83×10 ⁻⁴	1.40×10 ⁻⁴
Pyrene	6.40×10 ⁻⁵	1.98×10 ⁻⁴	1.52×10 ⁻⁴
Benz(a)anthracene	5.40×10 ⁻⁶	4.76×10 ⁻⁴	3.64×10 ⁻⁴
Chrysene	6.05×10 ⁻⁶	4.02×10 ⁻⁴	3.08×10 ⁻⁴
Benzo(a)pyrene	2.94×10 ⁻⁷	1.19×10 ⁻³	9.13×10 ⁻⁴
Benzo(b)fluoranthene	4.01×10 ⁻⁶	5.81×10 ⁻⁴	4.45×10 ⁻⁴
Benzo(k)fluoranthene	4.01×10 ⁻⁶	5.81×10 ⁻⁴	4.45×10 ⁻⁴
Benzo(g,h,i)perylene	0.0	3.23×10 ⁻³	2.47×10 ⁻³
Indeno(1,2,3,c,d)pyrene	0.0	1.21×10 ⁻³	9.28×10 ⁻⁴
Dibenzo(a,h)anthracene	0.0	2.79×10 ⁻⁵	2.13×10 ⁻⁵

Note: The zero values are estimated because of the zero-phase fractions from Table 2-42

Table 2-44 Description of the Database Tables pahGasRatio and pahParticleRatio

Field Description		Relevant Values
riciu	Description	ixcicvant values
polProcessID	Identifies the pollutant (1st two digits and Emissions Process (last two digits).	Pollutants are identified in the table above; Relevant polprocesses include: 18501 = "Naphthalene gas, running exhaust" 18502 = "Naphthalene gas, start exhaust"
fuelTypeID	Identifies broad classes of fuels, e.g., "gasoline." "diesel."	1 = "Gasoline" 2 = "Diesel" 3 = "CNG" 5 = "Ethanol"
modelYearGroupID	Identifies a set of model years covered by a specific value of atRatio.	1960-1970 1971-1977 1978-1995 1996-2006 2007-2060
atRatio	Average PAH/VOC emission ratio for a combination of process, fuel type, sourceType and modelYearGroup.	
meanBaseRateCV	"Coefficient of Variation of the Mean" or "relative standard error" of the meanBaseRate.	
dataSourceID	Indicates source data and methods used to estimate atRAtio.	

2.2.2 Vehicles Operating on Fuel Blends Containing 70-100 percent Ethanol

As noted above, flexible-fueled vehicles (FFVs) may operate on gasoline, or on "E85" fuel, which is a mix of gasoline with high ethanol levels (typically 70-85%). Hays et al. (2013)³⁴ reported speciated filter-collected semi-volatile organic compound (SVOC) measurements from three Tier 2 compliant vehicles tested using E0, E10 and E85 fuels. Reductions in total PAH between E0 and E85 in total measured filter-collected PAHs ranged between 22 percent and 93 percent depending on the temperature and phase of the LA92 cycle. They found that E85 significantly reduced the lighter PAHs, including naphthalene, fluorene, anthracene, phenanthrene, fluoranthene, pyrene, benzo(*a*)anthracene and chrysene. However, no significant effect was observed for the heavier PAHs, including benzo(*a*)pyrene, benzo(*k*)fluoranthene, benzo(*ghi*)perylene, and indeno(1,2,3-*cd*)pyrene.

Because Hays et al. (2013) reported only the filter-collected PAH emissions, and the results were conducted on a limited number of vehicles, we used the results to adjust the fleet-average PAH ratios derived from KCVES tested on E0 fuel. We reduced the VOC phase PAH ratios by 74 percent, assuming that (1) the annual average ethanol content of high ethanol fuels is 74 percent, and (2) the PAH in the gaseous phase are reduced proportionally to the gasoline content

reductions. The 74 percent reduction is within the range of reductions observed by Hays et al. (2013)³⁴ for total PAHs. Because Hays et al. (2013)³⁴ observed no significant decrease of the heavier PAHs for which MOVES assumes exist primarily in the particle-phase (Table 2-42), we assume the E85 particle PAH/OC fractions are the same as the E0-E20 fractions derived from KCVES. The resulting fractions are presented in Table 2-45.

Table 2-45 Toxic Fractions for PAH species for Vehicles Running on High-Ethanol Blends by Process

PAH species	Gaseous	Particulate Phase (PAH/OC _{2.5}	
	Phase (PAH/VOC)	Start	Running
Naphthalene	5.38×10 ⁻⁴	1.68×10 ⁻⁴	1.29×10 ⁻⁴
Acenaphthylene	4.71×10 ⁻⁵	5.01×10 ⁻⁵	3.83×10 ⁻⁵
Acenaphthene	1.04×10 ⁻⁵	0.0	0.0
Fluorene	2.10×10 ⁻⁵	0.0	0.0
Anthracene	8.70×10 ⁻⁶	5.19×10 ⁻⁵	3.97×10 ⁻⁵
Phenanthrene	5.57×10 ⁻⁵	1.81×10 ⁻⁴	1.39×10 ⁻⁴
Fluoranthene	1.45×10 ⁻⁵	1.83×10 ⁻⁴	1.40×10 ⁻⁴
Pyrene	1.66×10 ⁻⁵	1.98×10 ⁻⁴	1.52×10 ⁻⁴
Benz(a)anthracene	1.41×10 ⁻⁶	4.76×10 ⁻⁴	3.64×10 ⁻⁴
Chrysene	1.57×10 ⁻⁶	4.02×10 ⁻⁴	3.08×10 ⁻⁴
Benzo(a)pyrene	7.65×10 ⁻⁸	1.19×10 ⁻³	9.13×10 ⁻⁴
Benzo(b)fluoranthene	1.04×10 ⁻⁶	5.81×10 ⁻⁴	4.45×10 ⁻⁴
Benzo(k)fluoranthene	1.04×10 ⁻⁶	5.81×10 ⁻⁴	4.45×10 ⁻⁴
Benzo(ghi)perylene	0.0	3.23×10 ⁻³	2.47×10 ⁻³
Indeno(1,2,3,cd)pyrene	0.0	1.21×10 ⁻³	9.28×10 ⁻⁴
Dibenzo(ah)anthracene	0.0	2.79×10 ⁻⁵	2.13×10 ⁻⁵

Note: The zero values are estimated because of the zero-phase fractions from Table 2-42

As discussed in the MOVES Fuel Effects report²², the VOC and PM_{2.5} emission rates (g/mile or g/start) calculated using MOVES for vehicles fueled on E10 and E85 are quite similar, except for VOC emissions for pre-2001 MY vehicles. As such, the difference in the gas-phase PAH emission rates between E10 and E85 fueled-vehicles for the MY 2001 and later are driven by the differences in the toxic fractions shown in Table 2-45 (74% lower). Because the particle-phase PAH toxic ratios are the same between E85 and E10 vehicles, the particle-phase PAH emission rates for these vehicles from MOVES are roughly equivalent.

2.3 Metals

Emissions of metals in vehicle exhaust result from trace-level contamination of fuel and engine oil, as well as attrition from engine, exhaust system, and emission-control components. MOVES models two groups of metal emissions, 1) metals that are used for air quality modeling, and 2) metals that are included due to their known toxicity. The metals that are included for air quality modeling, which include metals such as iron, aluminum and calcium are discussed in the MOVES3 Speciation report.²¹ Emissions of these metals are estimated as fractions of PM_{2.5} emission rates.

This report covers seven metal species included due to their known toxicity, including five metals and three forms of mercury, as listed in Table 1-4. The toxic metal emissions are estimated using distance-specific emission rates (g/mile). Manganese is the only metal that is required for both air quality and toxicity purposes and is estimated using the g/mile approach. In the database, these rates are stored in the metalEmissionRate table, described in Table 2-47. Note that while the table contains a field for "fuel type," the emission rates listed in the table do not vary among fuel types.

Emission rates for magnesium and nickel were developed from the 99 vehicles sampled for chemical composition in the KCVES. The mean rates are calculated as weighted averages of metal measured on Bag 2 of the LA92, using weights designed to represent the onroad vehicle fleet. The use of Bag 2 emissions in the averaging helps ensure that the emission rates for these metals are consistent with the PM_{2.5} emission profile for running emissions discussed in the MOVES3 Speciation Report. These approaches were adopted because, while PM_{2.5} emissions are much lower during hot-stabilized running conditions, PM_{2.5} emissions are more enriched in metals during hot-stabilized running conditions than during start emissions. We compared the g/mi emission rates from Bag 2 to the average of the entire LA92; the difference in the Bag 2 emission rates from the average of the LA92 is 38 percent and -16 percent for manganese and nickel. Thus, in using Bag 2 emission rates for metal emission rates, the approach is both consistent with the PM_{2.5} speciation running emission profile and provides a likely upper limit (in the case of manganese) when compared to the cycle average.

Hexavalent chromium was estimated using data collected at U.S. EPA's National Vehicle Emissions Laboratory and analyzed at the Wisconsin State Laboratory of Hygiene at the University of Wisconsin-Madison. These data were collected on a single vehicle, a 2008 Chevrolet Impala flexible-fuel vehicle. At the time of the analysis, they were the only available data with direct measurement of hexavalent chromium from a highway vehicle. Development of a gasoline vehicle emission rate from these data is detailed in Appendix A.

Emission factors for arsenic were developed from data reported for tunnel tests.³⁶ These data were collected in two Milwaukee tunnels in 2000/2001, using inductively-coupled plasma mass spectrometry (ICP-MS). The emission rates are fleet-average, however the fleet had between 1.5% and 9.4% percentage of heavy-duty trucks. We use the same fleet-average emission rate for both gasoline and diesel vehicles.^h

Emission factors for mercury were obtained from a 2005 test program at EPA's National Exposure Research Laboratory (NERL). In this program mercury samples in raw exhaust were collected from 14 light-duty gasoline vehicles and two heavy-duty diesel vehicles. Documentation describing development of these emission factors can be found in Appendix B.

the arsenic may be coming from non-tailpipe or road dust sources.

h Schauer et a. (2006)³⁶ used a chemical mass balance (CMB) model to apportion metal emissions to diesel exhaust, combined gasoline tailpipe and tire wear, brake wear, and road dust sources. However, arsenic was not measured in significant quantities to conduct the CMD source apportionment. We used the total tunnel emission rates to represent arsenic exhaust emission rates in MOVES for both gasoline and heavy-duty vehicles, however, some of

Table 2-46 Metal Emission Rates for Gasoline Motor Vehicles

Pollutant	Emission Rate (g/mi)
Chromium, hexavalent (6+)	1.20×10 ⁻⁸
Manganese	1.33×10 ⁻⁶
Nickel	1.50×10 ⁻⁶
Mercury, Elemental (Gaseous Phase)	1.10×10 ⁻⁷
Mercury, Reactive (Gaseous Phase)	9.90×10 ⁻⁹
Mercury, Particulate Phase	4.00×10 ⁻¹⁰
Arsenic	2.30×10 ⁻⁶

Fleet-average metal emission rates were derived for vehicles running on gasoline and gasoline-ethanol blends. Since metal emissions can result from trace level contamination of fuel and engine oil, as well as wear on engine and exhaust aftertreatment components, the metal emission rates for E85-fueled vehicles were assumed to be the same as those used for gasoline vehicles (see Table 2-46).

Table 2-47 Description of the Database Table metalEmissionRate

Table 2-47 Description of the Database Table metalEmissionRate				
Field	Description	RelevantValues		
polProcessID	Identifies the pollutant (1st two digits and Emissions Process (last two digits).	Pollutants are identified in the table above; Relevant processes include: 1 = "Running Exhaust"		
fuelTypeID	Identifies broad classes of fuels, e.g., "gasoline." "diesel."	1 = "Gasoline" 2 = "Diesel" 5 = "Ethanol"		
sourceTypeID	Identifies vehicle types, classified by function	Motorcycles (11) Passenger Cars (21) Passenger Trucks (31) Light Commercial Trucks (32)		
modelYearGroupID	Identifies a set of model years covered by a specific value of atRatio.	1960-1970 1971-1977 1978-1995 1996-2006 2007-2060		
Units	Identifies units in which the meanBaseRate is expressed.	grams/mile		
meanBaseRate	Average emission rate for a combination of process, fuel type, sourceType and modelYearGroup.			
meanBaseRateCV	"Coefficient of Variation of the Mean" or "relative standard error" of the meanBaseRate.			
dataSourceID	Indicates source data and methods used to estimate atRAtio.			

2.4 Dioxins and Furans

The MOVES model estimates mass and distance-based emission rates for 17 dioxin and furan congeners (gram/mile). We discuss the derivation of dioxin and furan emission rates for vehicles operating on gasoline containing low ethanol percentages (Section 2.4.1) and high-ethanol percentages (Section 2.4.2).

2.4.1 Vehicles Operating on Fuel Blends Containing 0-15 Percent Ethanol

The emission rates for dioxins and furans were obtained from the tunnel study used in EPA's dioxin assessment.^{37,38} The emission rates from the tunnel study did not vary among fuel types and we applied theses rates to all gasoline vehicles in MOVES. In the absence of additional data, the fractions for more recently-manufactured vehicles were assumed to be the same as those for vehicles employing older technologies. Of course, this extrapolation from one set of technologies to another involves some degree of uncertainty. The rates are stored in the dioxinEmissionRate table, which is described in Table 2-49.

Table 2-48 Dioxin Emission Rates for Motor Vehicles Running on Gasoline Fuel Blends with 0-20 Percent Ethanol

Pollutant	mg/mi
2,3,7,8-Tetrachlorodibenzo- <i>p</i> -Dioxin (TCDD)	8.27×10 ⁻¹⁰
1,2,3,7,8-Pentachlorodibenzo-p-Dioxin	3.70×10 ⁻¹⁰
1,2,3,4,7,8-Hexachlorodibenzo-p-Dioxin	3.87×10 ⁻¹⁰
1,2,3,6,7,8-Hexachlorodibenzo-p-Dioxin	7.92×10 ⁻¹⁰
1,2,3,7,8,9-Hexachlorodibenzo-p-Dioxin	4.93×10 ⁻¹⁰
1,2,3,4,6,7,8-Heptachlorodibenzo-p-Dioxin	5.95×10 ⁻⁹
Octachlorodibenzo-p-dioxin	4.70×10 ⁻⁸
2,3,7,8-Tetrachlorodibenzofuran	2.76×10 ⁻⁹
1,2,3,7,8-Pentachlorodibenzofuran	1.32×10 ⁻⁹
2,3,4,7,8-Pentachlorodibenzofuran	9.68×10 ⁻¹⁰
1,2,3,4,7,8-Hexachlorodibenzofuran	1.09×10 ⁻⁹
1,2,3,6,7,8-Hexachlorodibenzofuran	1.16×10 ⁻⁹
1,2,3,7,8,9-Hexachlorodibenzofuran	3.17×10 ⁻¹⁰
2,3,4,6,7,8-Hexachlorodibenzofuran	1.36×10 ⁻⁹
1,2,3,4,6,7,8-Heptachlorodibenzofuran	1.21×10 ⁻⁸
1,2,3,4,7,8,9-Heptachlorodibenzofuran	3.87×10 ⁻¹⁰
Octachlorodibenzofuran	1.37×10 ⁻⁸

Table 2-49 Description of the Database Table DioxinEmissionRate

Table 2-45 Description of the Database Table Dioxine mission Rate			
Field	Description	RelevantValues	
polProcessID	Identifies the pollutant (1st two digits and Emissions Process (last two digits).	Pollutants are identified in the table above; Relevant processes include: 1 = "Running Exhaust"	
fuelTypeID	Identifies broad classes of fuels, e.g., "gasoline." "diesel."	1 = "Gasoline" 2 = "Diesel" 5 = "Ethanol"	
modelYearGroupID	Identifies a set of model years covered by a specific value of atRatio.	1960-2050 1960-2006 2007-2009 2010-2060	
Units	Identifies units in which the meanBaseRate is expressed.	grams/mile	
meanBaseRate	Average emission rate for a combination of process, fuel type, sourceType and modelYearGroup.		
meanBaseRateCV	"Coefficient of Variation of the Mean" or "relative standard error" of the meanBaseRate.		
dataSourceID	Indicates source data and methods used to estimate atRAtio.		

2.4.2 Vehicles Operating on Fuel Blends containing 70-100 percent Ethanol

No emissions data exist for dioxin and furan emissions from vehicles running on E85 or E70. Thus, dioxin emission factors for E85 and E70 were estimated by multiplying fractions for vehicles running on E0 fuels (Table 2-48) by the fraction of gasoline in the fuel, assuming no emission of dioxins or furans resulting from the combustion of ethanol. Resulting ratios are given in Table 2-50.

Table 2-50 Emission Factors for Dioxins and Furans, for Vehicles Operating on High-Ethanol Blends

Congener	Emission rate
2 2 7 9 T-41111	(mg/mile) 2.15×10 ⁻¹⁰
2,3,7,8-Tetrachlorodibenzo-p-dioxin	
1,2,3,7,8-Pentachlorodibenzo-p-Dioxin	9.61×10 ⁻¹¹
1,2,3,4,7,8-Hexachlorodibenzo-p-Dioxin	1.01×10 ⁻¹⁰
1,2,3,6,7,8-Hexachlorodibenzo-p-Dioxin	2.06×10 ⁻¹⁰
1,2,3,7,8,9-Hexachlorodibenzo-p-Dioxin	1.28×10 ⁻¹⁰
1,2,3,4,6,7,8-Heptachlorodibenzo-p-Dioxin	1.55×10 ⁻⁹
Octachlorodibenzo-p-dioxin	1.22×10 ⁻⁸
2,3,7,8-Tetrachlorodibenzofuran	7.19×10 ⁻¹⁰
1,2,3,7,8-Pentachlorodibenzofuran	3.43×10 ⁻¹⁰
2,3,4,7,8-Pentachlorodibenzofuran	2.52×10 ⁻¹⁰
1,2,3,4,7,8-Hexachlorodibenzofuran	2.84×10 ⁻¹⁰
1,2,3,6,7,8-Hexachlorodibenzofuran	3.02×10 ⁻¹⁰
1,2,3,7,8,9-Hexachlorodibenzofuran	8.24×10 ⁻¹¹
2,3,4,6,7,8-Hexachlorodibenzofuran	3.52×10 ⁻¹⁰
1,2,3,4,6,7,8-Heptachlorodibenzofuran	3.16×10 ⁻⁹
1,2,3,4,7,8,9-Heptachlorodibenzofuran	1.01×10 ⁻¹⁰
Octachlorodibenzofuran	3.57×10 ⁻⁹

3 Diesel Exhaust

Toxic fractions, dioxin and metal emission rates were developed for exhaust emissions from heavy-duty diesel vehicles and applied to all diesel vehicle categories. The development of inputs for diesel vehicles are defined as "pre-2007", "model year 2007 through 2009", and "2010 and later" based on technology and emissions standards for heavy-duty vehicles. These distinctions are made because emission controls on 2007 through 2009 and 2010 and later engines differ and have a substantial effect on composition of emissions. The pre-2007 diesel toxic fractions for VOCs and PAHs are applied to auxiliary power unit exhaust for all model year vehicles until 2024, because auxiliary power units are not subject to the same stringency of control as highway engines. There are no separate emission ratios or factors for diesel engines running on biodiesel fuels or synthetic diesel fuels, due to limited data. Biodiesel vehicles use the same toxic ratios and factors as regular diesel. Due to a lack of applicable data, the toxic emission data are based on heavy-duty testing but are applied to light-duty diesel with the same model year distinctions (pre-2007, 2007 to 2009, and 2010 and later).

3.1 Volatile Organic Compounds

The composition of VOC emissions for heavy-duty diesel engines lacking the advanced control technologies applied in more recently-manufactured vehicles differs substantially from earlier technologies. Thus, we developed one set of toxic fractions for pre-2007 diesel engines and another set for engines manufactured between 2007 to 2009, and 2010 and later. For each model year group, the toxic fractions were developed from test cycles that included both running and start exhaust emissions and the same toxic fractions are applied to both emission processes in MOVES.

3.1.1 Pre-2007 Diesel Engines

To estimate toxic fractions of VOC for vehicles in the pre-2007 model-year group, EPA relied on a database compiled for the Coordinating Research Council and the National Renewable Energy Laboratory (NREL) (CRC E-75).³⁹ This database was developed from a literature survey and compiled data collected in 13 different studies. The studies included were conducted in various countries and included heavy-duty and light-duty engines, a variety of diesel and biodiesel fuels, and a number of different operating modes and cycles.

For 2,2,4-trimethylpentane, hexane, propionaldehyde, and toluene, toxic fractions of VOC were developed by Sierra Research. Their analysis of CRC E-75 data is described in detail in the technical report. ^{45,39} Data from tests using non-conventional diesel fuel (Fischer-Tropsch, biodiesel, ethanol-diesel blends, emulsified fuel, European blends, and other obvious research fuels) were excluded, as were data from light-duty engines. A single set of toxic fractions were developed from the measurements on different test cycles, with no differentiation between cycles with start emissions and running emissions only. ³⁹ The fractions are provided in Table 3-1. Toxic fractions for other compounds in Table 3-1 were developed by EPA from the E-75 database. We relied on data collected in the United States from heavy-duty diesel engines running on conventional diesel fuels, collected on test-cycles representative of real-world operation. Some

studies reported results on a distance-specific basis (g/mi) whereas others reported results on a brake-specific basis (g/hp-hr). For both subsets of data, we calculated mean emissions for each toxic and for VOC, and then calculated mean fractions for each reporting basis. We then calculated an overall mean fraction using the respective sample sizes to weight the two fractions.

Table 3-1 Toxic Fractions of VOC for Pre-2007 Diesel Engines

Pollutant	Toxic fraction
1,3-Butadiene	0.00292
2,2,4-Trimethylpentane	0.00180
Acetaldehyde	0.03555
Acrolein	0.00662
Benzene	0.00783
Ethanol	0
Ethyl Benzene	0.00266
Formaldehyde	0.07823
n-Hexane	0.00197
Propionaldehyde	0.00468
Styrene	0.00131
Toluene	0.00433
Xylenes	0.00378

Since extended idle emissions associated with auxiliary power units (APUs) are not subject to 2007 standards, toxic to VOC ratios for pre-2007 diesel engines were used for the APU VOC toxic emission rates for all model years until 2024. However, we anticipate that APU standards promulgated as part of the Phase 2 greenhouse gas regulation for medium and heavy-duty engines will result in use of diesel particulate filters in 2024; thus we are using 2007-2009 diesel toxic emissions data for those units.

3.1.2 2007+ Diesel Engines

For heavy-duty diesel engines manufactured in 2007 and later, advanced emission controls change the composition of VOCs. For these engines, we relied on speciated emissions data from the Advanced Collaborative Emissions Study (ACES), directed by the Health Effects Institute and Coordinating Research Council, with participation from a range of government and privatesector sponsors. 42, 43 ACES was conducted in two phases, with the first focusing on engines complying with 2009 standards and the second focusing on engines meeting 2010 standards. Engines tested in ACES that met the 2007 and 2010 standards had different emission control systems which had significant impact on composition of emissions. Whereas 2007-compliant engines added diesel particulate filters, 2010-compliant engines also had urea-based selective catalytic reduction (SCR) catalysts and ammonia oxidation catalysts. In ACES Phase 1, detailed emissions measurements were performed on four engines, while in ACES Phase 2 measurements were performed on three engines. In both test programs, vehicles were operated on low-sulfur diesel fuel over several test cycles, with new engine and aftertreatment systems. We made use of data from the 16-hour transient cycle which is composed of FTP and CARB 5-Mode cycles, developed specifically to gain sufficient mass of toxics emitted at low concentrations, and to capture diesel particulate filter regeneration events. The ACES measurements for the selected VOC emissions were background-corrected using background dilution air. 42 Toxic fractions of

VOC calculated from the ACES data are provided in Table 3-2. Because VOC emissions are so low in the advanced technology diesels with new engine and aftertreatment systems, there is considerable measurement uncertainty for gaseous air toxics. Also, while VOC measurements included several alcohol species, including ethanol, these measurements were likely due to sample contamination and were not included in the speciation profiles used to develop toxic fractions, (See discussion in Appendix D of the Speciation Report²¹) and the ethanol toxic fraction is set to zero (Table 3-2).

Table 3-2 Toxic Fractions of VOC for 2007 and Later Diesel Vehicles

	Toxic Fraction		
Pollutant	2007-2009	2010 and later	
1,3-Butadiene	0.0008	0.0000	
2,2,4-Trimethylpentane	0.0078	0.0045	
Acetaldehyde	0.0693	0.0417	
Acrolein	0.0100	0.0036	
Benzene	0.0129	0.0000	
Ethanol	0	0	
Ethyl Benzene	0.0063	0.0112	
Formaldehyde	0.2174	0.0266	
N-Hexane	0.0054	0.0009	
Propionaldehyde	0.0031	0.0029	
Styrene	0.0000	0.0000	
Toluene	0.0300	0.0183	
Xylene(s)	0.0380	0.0848	

3.2 Polycyclic Aromatic Hydrocarbons

PAH emissions from diesel vehicles can be formed by combustion as well as post-combustion in the engine and aftertreatment systems.⁴⁷ We have developed separate PAH emission rates for pre-2007 diesel, 2007-2009 diesel, and 2010+ diesel to represent the different combustion and aftertreatment strategies as discussed in the following subsections.

As with gasoline emissions, PAH mass emissions from diesel engines were apportioned into gaseous and particulate phases, using a single set of allocation factors for all temperature conditions. The partitioning factors for diesel PAHs were developed using estimates from EPA's SPECIATEdatabase⁴⁴ and information on compounds' physical and chemical properties. ⁴⁵ The allocations from SPECIATE were based on exhaust measured from two medium-duty diesel trucks tested in 1996 with low mileage. ⁴⁶ The phase-partitioning factors are shown in Table 3-3. Compared to the partitioning for gasoline (Table 2-42), the fraction of PAH in the particulate phase is higher for diesel emissions, which is consistent with the higher concentrations of particles in diesel exhaust. It should be noted that the data used represent gas-particle phase partitioning in the sampled diluted exhaust, which is not fully representative of partitioning in the atmosphere.

Emissions of PAH in the gaseous and particulate phases were estimated as fractions of total VOC and OC_{2.5}, respectively. We used the phase-partitioning factors in Table 3-3 for both pre-2007 and 2007+ diesel engines. We note that we introduce additional uncertainty when we apply

the phase-partitioning derived from two pre-2007 technology vehicles to the 2007+ diesel engines

Table 3-3 Phase-Partition Fractions for Emissions of Polycyclic Aromatic Hydrocarbons from Diesel Engines

PAH species	Molar Mass (g/mol)	Phase Fraction	
		Gaseous	Particulate
Naphthalene	128	1.0	0.0
Acenaphthylene	152	1.0	0.0
Acenapthene	154	1.0	0.0
Fluorene	166	0.785	0.215
Anthracene	178	0.534	0.466
Phenanthrene	178	0.665	0.335
Fluoranthene	202	0.484	0.516
Pyrene	202	0.448	0.552
Benz(a)anthracene	228	0.277	0.723
Chrysene	228	0.177	0.823
Benzo(a)pyrene	252	0.0	1.0
Benzo(b)fluoranthene	252	0.0	1.0
Benzo(k)fluoranthene	252	0.0	1.0
Benzo(ghi)perylene	276	0.227	0.773
Indeno(1,2,3-cd)pyrene	276	0.0	1.0
Dibenzo(ah)anthracene	278	0.0	1.0

3.2.1 Pre-2007 Diesel Engines

PAH fractions for pre-2007 diesel engines were calculated using results from the E-75 database. A single set of PAH/VOC fractions were derived for the E-75 database that represent both start and running exhaust. For the particulate phase, a single fraction was first calculated with respect to total PM_{2.5}, and then converted to a fraction of total OC_{2.5} using estimates of OC as a fraction of total PM_{2.5}. Note that the OC/PM fractions differed by emissions process, thus for MOVES we obtained separate PAH/OC_{2.5} fractions for start, running and extended-idle emissions.

In estimating fractions, we relied on data collected in the United States on heavy-duty diesel engines running on conventional diesel fuels, measured on test-cycles representative of real-world operation. It should be noted that for some compounds, substantially more data were available than for others; thus, the level of confidence in emission rates varies among individual compounds. For instance, while data from 66 tests were available for acenaphthene, data from only two tests were available for dibenz(*ah*)anthracene. Table 3-4 shows fractions for PAH emissions relative to OC and VOC, by emissions process.

Table 3-4 Toxic Fractions for PAH Species, by Phase and Process, for Pre-2007 Diesel Vehicles

РАН	Gaseous	Partic	ulate Phase (PAH/OC _{2.5})
	Phase (PAH/VOC)	Start/Idle	Running	Extended Idle
Naphthalene	9.05×10 ⁻³	0.0	0.0	0.0
Acenaphthylene	5.01×10 ⁻⁴	0.0	0.0	0.0
Acenaphthene	2.98×10 ⁻⁴	0.0	0.0	0.0
Fluorene	4.85×10 ⁻⁴	2.80×10 ⁻⁴	8.49×10 ⁻⁴	2.54×10 ⁻⁴
Anthracene	2.35×10 ⁻⁴	1.63×10 ⁻⁴	4.94×10 ⁻⁴	1.48×10 ⁻⁴
Phenanthrene	7.08×10 ⁻⁴	6.44×10 ⁻⁴	1.96×10 ⁻³	5.86×10 ⁻⁴
Fluoranthene	3.55×10 ⁻⁴	6.24×10 ⁻⁴	1.90×10 ⁻³	5.68×10 ⁻⁴
Pyrene	4.27×10 ⁻⁴	9.02×10 ⁻⁴	2.74×10 ⁻³	8.21×10 ⁻⁴
Benzo(a)anthracene	4.36×10 ⁻⁵	3.23×10 ⁻⁴	9.81×10 ⁻⁴	2.94×10 ⁻⁴
Chrysene	1.70×10 ⁻⁵	2.04×10 ⁻⁴	6.20×10 ⁻⁴	1.86×10 ⁻⁴
Benzo(a)pyrene	0.0	1.21×10 ⁻⁴	3.69×10 ⁻⁴	1.10×10 ⁻⁴
Benzo(b)fluoranthene	0.0	3.60×10 ⁻⁵	1.10×10 ⁻⁴	3.28×10 ⁻⁵
Benzo(k)fluoranthene	0.0	5.08×10 ⁻⁶	1.50×10 ⁻⁵	4.62×10 ⁻⁶
Benzo(ghi)perylene	8.3×10 ⁻⁷	5.78×10 ⁻⁶	1.80×10 ⁻⁵	5.26×10 ⁻⁶
Indeno(1,2,3-cd)pyrene	0.0	9.24×10 ⁻⁶	2.81×10 ⁻⁵	8.41×10 ⁻⁶
Dibenz(ah)anthracene	0.0	4.85×10 ⁻⁶	1.50×10 ⁻⁵	4.41×10 ⁻⁶

Note: The zero values are estimated because of the zero-phase fractions from Table 3-3

The PAH Toxic fractions in Table 3-4 are applied to exhaust emission for 2006 and earlier model year diesel vehicles in MOVES. The extended idle toxic fractions are applied to auxiliary power unit (APUs) exhaust for all model year vehicles in MOVES prior to 2024, because the APUs are not subject to the same control as exhaust from the highway engines. However, as previously discussed, we are using 2007-2009 diesel toxic emissions data for 2024 and later units.

3.2.2 2007+ Diesel Engines

Advanced emission controls used in 2007 and later model year diesel including diesel vehicles particulate filters, diesel oxidation catalysts, and selective catalytic reduction (SCR) catalysts and ammonia oxidation catalysts can reduce or facilitate the formation of individual PAHs. PAHs are destroyed through oxidation by diesel oxidation catalysts, catalyzed-diesel particulate filters and NO₂ present in the aftertreatment that is produced by the diesel particulate filter but controlled by the SCR^{47,48} Modern aftertreatment systems can also facilitate the formation of individual PAH emissions. For example, Liu et al. 2015⁴⁷ measured an increase in nitro-PAHs emissions in some tests configurations of a nonroad diesel engine with an oxidation catalyst + SCR aftertreatment system compared to a baseline case with no diesel aftertreatment systemⁱ.

ⁱ Nitro-PAHs are not currently modeled in MOVES

For these engines, we relied on speciated emissions data from the ACES studies^{42,43} The PAH emissions measured in the ACES study were uncorrected for background concentrations.⁴² The 16-hour drive cycle used in the ACES studies comprises multiple driving modes and is used to represent both start and running exhaust in MOVES. Toxic fractions applicable to these engines are shown in Table 3-5, in which the fractions are differentiated by phase (gas or particulate) but not by emissions process (start and running). We used the same phase fractions presented in Table 3-3. For the particulate phase, a single fraction is provided for all processes (similar to VOC) because the OC/PM fraction in MOVES for 2007+ diesel is a single fraction for both start and running emission processes.²¹ The 2007 and later engines have lower gaseous PAH fractions than the pre-2007 engines with the exception of naphthalene, and phenanthrene, and most of the particulate phase PAHs. The 2010 and later engines emissions technologies facilitate further reductions in PAHs and have significantly lower fractions of all gaseous and particle-phase PAH emission rates than the 2007-2009 emission rates, with the exception of benzo(a)anthracene.

Table 3-5 Toxic Fractions for Polycyclic Aromatic Compounds, by Phase, for 2007 and Later Diesel Vehicles

Table 3-5 Toxic Fractions	2007-2009		2010 and later	
РАН	Gaseous Phase (PAH/VOC)	Particulate Phase (PAH/OC2.5)	Gaseous Phase (PAH/VOC)	Particulate Phase (PAH/OC2.5)
Naphthalene	1.63×10 ⁻²	0.0	5.84×10 ⁻⁴	1.35×10 ⁻⁵
Acenaphthylene	8.53×10 ⁻⁵	0.0	1.49×10 ⁻⁵	1.29×10 ⁻⁶
Acenaphthene	5.26×10 ⁻⁵	0.0	1.56×10 ⁻⁵	0
Fluorene	1.96×10 ⁻⁴	2.41×10 ⁻⁴	3.35×10 ⁻⁵	0
Anthracene	3.04×10 ⁻⁵	1.19×10 ⁻⁴	6.47×10 ⁻⁶	3.19×10 ⁻⁶
Phenanthrene	8.51×10 ⁻⁴	1.92×10 ⁻³	9.62×10 ⁻⁵	2.61×10 ⁻⁵
Fluoranthene	4.57×10 ⁻⁵	2.18×10 ⁻⁴	6.41×10 ⁻⁶	6.684×10 ⁻⁶
Pyrene	3.79×10 ⁻⁵	2.09×10 ⁻⁴	4.72×10 ⁻⁶	4.67×10 ⁻⁶
Benzo(a)anthracene	3.00×10 ⁻⁷	3.58×10 ⁻⁶	6.92×10 ⁻⁷	1.942×10 ⁻⁵
Chrysene	5.00×10 ⁻⁷	1.12×10 ⁻⁵	2.51×10 ⁻⁷	5.32×10 ⁻⁶
Benzo(a)pyrene	0.0	1.48×10 ⁻⁵	0.0	0.0
Benzo(b)fluoranthene	0.0	6.27×10 ⁻⁶	0.0	0.0
Benzo(k)fluoranthene	0.0	6.27×10 ⁻⁶	0.0	0.0
Benzo(ghi)perylene	2.00×10 ⁻⁷	8.96×10 ⁻⁷	0.0	0.0
Indeno(1,2,3-cd)pyrene	0.0	2.24×10 ⁻⁶	0.0	0.0
Dibenz(a,h)anthracene	0.0	4.48×10 ⁻⁶	0.0	0.0

Note: The zero values are estimated because of the zero-phase fractions from Table 3-3

3.3 Metals

Emission rates for selected metals representing pre-2007 heavy-duty diesel engines were based on data from the CRC E-75 program, with the exception of rates for hexavalent chromium, mercury and arsenic. The hexavalent chromium emission rate was obtained by multiplying the diesel vehicle emission rate by the ratio of hexavalent chromium to total chromium estimated

from gasoline exhaust (0.29%). The total chromium estimates for diesel vehicles came from the CRC E-75. More details are provided in Appendix A. The pre-2007 diesel emission rate for arsenic is a fleet-average value that is also used for gasoline vehicles, obtained from a fleet-average tunnel study (see Table 2-46). It does not vary with emission control technology. The mercury emission rate for pre-2007 diesels is calculated from emission tests conducted on two heavy-duty diesel vehicles, as documented in Appendix B. Table 3-6 provides metal emission factors for all diesel vehicles.

Table 3-6 Emission Rates for Selected Metals for Diesel Vehicles

Pollutant	Emission Rate for	Emission Rate for	Emission Rate for
	1960-2006 (g/mi)	2007-2009 (g/mi)	2010 and later (g/mi)
Chromium VI	2.00x10 ⁻⁸	5.93x10 ⁻⁹	2.16x10 ⁻⁹
Manganese	8.00×10 ⁻⁶	6.82×10 ⁻⁷	2.00x10 ⁻⁷
Nickel	1.40×10 ⁻⁵	6.92×10 ⁻⁷	2.63x10 ⁻⁷
Mercury, Elemental Gaseous	6.20×10 ⁻⁹	6.20×10 ⁻⁹	6.20x10 ⁻⁹
Phase			
Mercury, Reactive Gaseous	3.20×10 ⁻⁹	3.20×10 ⁻⁹	3.20x10 ⁻⁹
Phase			
Mercury, Particulate Phase	1.60×10 ⁻⁹	1.60×10 ⁻⁹	1.60x10 ⁻⁹
Arsenic	2.30×10 ⁻⁶	2.30×10 ⁻⁶	2.30x10 ⁻⁶

Emissions rates for manganese and nickel representing diesel engines manufactured in 2007-2009 were developed from the ACES Phase 1 program⁴², and 2010 and later were developed from the ACES Phase 2 program.⁴³ The ACES reported g/hp-hr rates from the 16-hour cycle were used, and converted to g/mile using the equation on page 31 of the ACES Phase 1 final report,⁴² and the measured brake-specific fuel consumption and an assumed heavy-duty fuel economy of 6 miles per gallon. The ACES metal emission rates were uncorrected for background concentrations.⁴² The emission rate for arsenic is identical to the emission rate used for gasoline vehicles and pre-2007 diesels (Table 2-46). The emission rates for mercury are the same as those derived for pre-2007 diesel engines, as discussed in Appendix B. The hexavalent chromium emission rates for pre-2007, 2007 to 2009, and 2010 and later diesel engines were obtained by multiplying the gasoline vehicle emission rate by the ratio of total chromium from diesel and gasoline engines. The total chromium estimates came from the previously cited KCVES and ACES test programs, respectively. More details are provided in Appendix A.

3.4 Dioxins and Furans

To represent emissions of dioxins and furans from pre-2007 heavy-duty diesel engines, emissions rates for 17 congeners were calculated from the results of an EPA diesel dioxin/furan study of legacy engines. In this study, dioxin emissions from three heavy-duty engines manufactured prior to 1994 were measured. These engines included a 1985 GM 6.2 L, a 1987 Detroit Diesel 6V92 and 1993 Cummins L10. The emission factors in mg/mi are shown in Table 3-7. Since these engines are older than most of the pre-2007 fleet, dioxin emissions for pre-2007 engines may be overestimated.

Table 3-7 Emission Rates for Dioxin/Furan Congeners for Diesel Vehicles (mg/mi)

Congener	1970-2006	2007 - 2009	2010 and later
2,3,7,8-Tetrachlorodibenzo- <i>p</i> -dioxin (TCDD)	2.23×10^{-10}	ND	ND
1,2,3,7,8-Pentachlorodibenzo-p-dioxin	ND	ND	ND
1,2,3,4,7,8-Hexachlorodibenzo- <i>p</i> -dioxin	ND	ND	ND
1,2,3,6,7,8-Hexachlorodibenzo- <i>p</i> -dioxin	1.03×10^{-10}	ND	ND
1,2,3,7,8,9-Hexachlorodibenzo- <i>p</i> -dioxin	4.78×10^{-10}	4.11 × 10 ⁻¹¹	ND
1,2,3,4,6,7,8-Heptachlorodibenzo-p-dioxin	4.18×10^{-9}	2.58×10 ⁻¹⁰	1.05×10 ⁻⁹
Octachlorodibenzo-p-dioxin	1.61×10^{-8}	9.30×10 ⁻¹⁰	6.98×10 ⁻⁹
2,3,7,8-Tetrachlorodibenzofuran	6.50×10^{-9}	ND	5.09×10 ⁻¹¹
1,2,3,7,8-Pentachlorodibenzofuran	1.39×10^{-9}	ND	1.07×10 ⁻¹⁰
2,3,4,7,8-Pentachlorodibenzofuran	2.23×10^{-9}	6.30×10 ⁻¹¹	3.24×10 ⁻¹⁰
1,2,3,4,7,8-Hexachlorodibenzofuran	8.02×10^{-10}	ND	2.20×10 ⁻¹⁰
1,2,3,6,7,8-Hexachlorodibenzofuran	4.24×10^{-10}	ND	2.43×10 ⁻¹⁰
1,2,3,7,8,9-Hexachlorodibenzofuran	ND	ND	ND
2,3,4,6,7,8-Hexachlorodibenzofuran	3.03×10^{-10}	ND	1.80×10 ⁻¹⁰
1,2,3,4,6,7,8-Heptachlorodibenzofuran	2.16×10^{-9}	3.00×10 ⁻¹⁰	9.94×10 ⁻¹⁰
1,2,3,4,7,8,9-Heptachlorodibenzofuran	ND	ND	5.81×10 ⁻¹¹
Octachlorodibenzofuran	1.85×10^{-9}	7.06×10 ⁻¹⁰	1.74×10 ⁻⁹

Note: ND = non-detected, fractions set to 0.

The data used to calculate the emission rates for engines manufactured between 2007 and 2009 were obtained from the EPA diesel dioxin study of 2007 and later engines. ⁴⁹ The results represent measurements during transient tests conducted on a MY2008 Cummins ISB engine over 48 replicates on the FTP cycle in a 1:23 cold:hot start ratio, combined with several emission-control technologies. To represent emissions from engines manufactured between 2007-2009, the results for the diesel oxidation-catalyst plus catalyzed diesel particulate filter were used. For engines manufactured in 2010 and later, the results for the diesel oxidation catalyst plus catalyzed diesel particulate-filter coupled with flow-through copper zeolite selective catalytic reduction and urea and ammonia slip catalyst were used. The 2007-2009 and 2010 and later emission rates are presented in Table 3-7.

4 Compressed Natural Gas (CNG) Exhaust

MOVES3 estimates emissions of toxics from heavy-duty vehicles fueled by compressed natural gas. This section describes the development of toxic emission inputs for this class of vehicles.

4.1 Volatile Organic Compounds

We used speciated hydrocarbon measurements from work sponsored by the California Air Resources Board. ⁵⁰ These measurements were taken on a 2000 MY Detroit Diesel Series 50G engine with and without an oxidation catalyst, measured on the Central Business District (CBD) driving cycle. As discussed in the MOVES3 Speciation report²¹, we used the uncontrolled results to represent speciation from pre-2002 CNG heavy-duty vehicles, and the results with oxidation-catalyst to represent 2002-and-later model year vehicles. The use of the CBD cycle is also consistent with the results used for criteria-pollutant emissions.

The toxic fractions of VOC derived from this set of measurements are displayed in Table 4-1. The total VOC emission rates with oxidation catalyst are reduced by 70 percent from pre-2002 levels. As shown in the table, formaldehyde emissions are preferentially reduced by the oxidation catalyst. Formaldehyde contributes over 50 percent of the VOC emissions for the uncontrolled CNG bus, but only 16.2 percent of the VOC emissions for the CNG bus equipped with an oxidation catalyst. The MOVES toxics not measured in this study are assumed to be negligible and are modeled as 0. The toxic fractions for VOC were applied to both start and running exhaust emissions in MOVES.

Table 4-1 Toxic Fractions of VOC for CNG Heavy-Duty Vehicles

	No control (pre-2002)	With oxidation catalyst (2002+)
1,3 Butadiene	0.000234	ND
Benzene	0.00135	0.00253
Toluene	0.000691	0.00786
Ethylbenzene	0.0000841	0.00131
Xylenes	0.000823	0.00634
Formaldehyde	0.517	0.162
Acetaldehyde	0.0305	0.138
Acrolein	0.00235	ND
Propionaldehyde	0.0153	ND

Note: ND = non detect, fractions set to 0.

4.2 Polycyclic Aromatic Hydrocarbons

The PAH toxic fractions for compressed natural gas are derived from tests on a model year 2000 DDC Series 50G engine on a New Flyer CNG transit bus tested by the California Air Resources Board (CARB).⁵¹ This engine had no catalyst, but the emission fractions are used to represent both catalyst and non-catalyst engines. Emissions were measured in two stages (the bus was re-

tested after 3 months of service in the Los Angeles County Metropolitan Transit Authority). The PAH emissions were measured in the semi-volatile phase using PUF-XAD and measured in the particulate phase on Teflon-coated glass-fiber filters. VOC emissions are derived from the NMHC and speciated hydrocarbon emissions. The OC emissions rates were provided to EPA by CARB. We estimated the volatile PAH emissions by calculating PAH/VOC fractions from the PUF-XAD measurements, and particle-phase PAH/OC fractions using the filter-based measurements for both stages of the study. For use in MOVES, we averaged the ratios estimated from both stages of the testing. The average ratios are displayed in Table 4-2. The PAH fractions were applied to both start and running exhaust emissions in MOVES.

Table 4-2 PAH Fractions of Volatile Organic Carbon (Volatile PAHs), and of Organic Carbon (Particle-Phase for CNG Heavy-Duty Vehicles

Thase for CNG Heavy-Duty Vehicles			
	Gaseous Phase (PAH/VOC)	Particulate Phase (PAH/OC)	
Compound	fraction	fraction	
Naphthalene	9.55×10 ⁻⁶	2.11×10 ⁻⁵	
Acenaphthylene	4.23×10 ⁻⁶	ND	
Acenaphthene	1.24×10 ⁻⁶	1.89×10 ⁻⁵	
Fluorene	2.99×10 ⁻⁶	3.30×10 ⁻⁵	
Anthracene	1.16×10 ⁻⁶	1.64×10 ⁻⁶	
Phenanthrene	8.36×10 ⁻⁶	2.04×10 ⁻⁵	
Fluoranthene	1.94×10 ⁻⁶	2.87×10 ⁻⁵	
Pyrene	3.74×10 ⁻⁶	5.35×10 ⁻⁵	
Benz(a)anthracene	1.68×10 ⁻⁷	9.39×10 ⁻⁶	
Chrysene/triphenylene	2.44×10 ⁻⁷	1.91×10 ⁻⁵	
Benzo(a)pyrene	ND	ND	
Benzo(b)fluoranthene	ND	ND	
Benzo(k)fluoranthene	ND	ND	
Indeno(1,2,3-cd)pyrene	ND	ND	
Benzo(ghi)perylene	ND	5.50×10 ⁻⁶	
Dibenz(ah)anthracene	ND	ND	

Note: ND = non detect, fractions set to 0.

4.3 Metals

We used the nickel emission rates reported from an uncontrolled 2000 MY DDC Series 50G engine.⁵² We used the uncontrolled engine to be consistent with the PM_{2.5} speciation profile. The hexavalent chromium emission rate was obtained by multiplying the total chromium from the DDC Series 50G CNG engine by the ratio of hexavalent chromium to total chromium from gasoline emissions (0.29%). More details are provided in Appendix A.

Results for the other metals predicted by MOVES were not available in the published literature. We used the same emission rates as for gasoline vehicles, because CNG heavy-duty vehicles are

largely spark ignition technology, with modern CNG vehicles having 3-way catalysts like light-duty gasoline vehicles. The rates are presented in Table 4-3.

Table 4-3 Metal Emission Rates and Sources used for CNG Vehicles

Pollutant	Emission Rate (g/mi)	Source
		University of Wisconsin
		(Appendix A) and Okamoto
Chromium 6+	2.1×10 ⁻¹⁰	et al. (2006)
Manganese	1.33×10 ⁻⁶	Same as gasoline
Nickel	1.00×10 ⁻⁸	Okamoto et al. (2006)
Elemental Gas Phase Hg	1.10×10 ⁻⁷	Same as gasoline
Reactive Gas Phase Hg	9.90×10 ⁻⁹	Same as gasoline
Particulate Hg	4.00×10 ⁻¹⁰	Same as gasoline
Arsenic	2.30×10 ⁻⁶	Same as gasoline

4.4 Dioxins and Furans

No published dioxin and furan emission rates for CNG vehicles were available. We are using the dioxin emission rates for gasoline reported in Table 2-48, because they both utilize spark ignition engines.

5 Evaporative and Refueling Emissions

Emissions of toxics emitted through evaporation of unburned fuel are estimated as fractions of total evaporative VOC. MOVES estimates toxic emission ratios for each evaporative process from gasoline vehicles (including gasoline-ethanol blends) and for refueling emissions from diesel vehicles. Currently, MOVES does not estimate evaporative emissions (e.g., refueling natural gas leaks) from CNG vehicles as discussed in the evaporative emission report.⁵³ This section documents the source of the toxic ratios used for evaporative emissions from gasoline and diesel vehicles.

5.1 Gasoline Vehicles

The derivation of the toxic fractions for vapor venting, fuel leaks and refueling emission processes are documented in Section 5.1.1 and for permeation in Section 5.1.2.

5.1.1 Vapor Venting, Fuel Leaks, and Refueling Emission Processes

MOVES estimates evaporative emissions from gasoline vehicles using toxic fractions that pertain to evaporative emission processes. The toxic fractions for some compounds are estimated as complex fractions based on fuel properties such as oxygenate content and vapor pressure. For other compounds, simple fractions are estimated. For the compounds modeled, fraction types and data sources are summarized in Table 5-1.

Expressions used to generate complex fractions were adapted from those used in MOBILE6.2.⁵⁴ These equations were adapted to compensate for a lack of data from newer vehicles collected in the context of appropriate experimental designs. However, as the conceptual basis for modeling evaporative emissions differs between MOBILE6 and MOVES, the equations are applied to the emission processes considered most closely analogous. Thus, equations for hot soak in MOBILE6.2 are used for vapor venting and refueling vapor loss, and equations for running loss are used for fuel leaks and refueling spillage loss. The equations are applied for fuels containing up to 20 percent ethanol and are presented in Table 5-2. We do not include naphthalene emissions from evaporative processes in MOVES since it is inconsistently measured in detectable quantities in evaporative emission testing. MOVES has fields for evaporative naphthalene, but all values in the model are zero.

Simple fractions for other air toxics in evaporative non-permeation emissions were obtained from profiles developed for EPA by Environ Corporation, using data from the Auto/Oil program conducted in the early 1990's. 55 The fractions for these compounds are the same for all pollutant processes (except permeation) and are presented in Table 5-3.

The ratios for 10 percent ethanol are used for all fuels with greater than or equal to 5 percent ethanol and less than 12 percent.

For vehicles operating on fuels containing 15 percent ethanol (E15), no data describing evaporative emissions are available. For the vapor-venting and spillage emission processes,

emission rates calculated from E15 and E10 fuel speciation data from the EPAct Program were used to adjust the E10 evaporative emissions speciation. ¹⁵ Resulting toxic fractions are provided in Table 5-3.

For vehicles containing 20 percent ethanol, toxic fractions were developed for fuel speciation profiles created from data collected in the EPAct program. Average fractions by weight were calculated as a composite of data from the seven E20 blends included in the fuel matrix. Resulting fractions are shown in Table 5-3. Note, MOVES versions after MOVES2014a do not estimate emissions for vehicles using E20 fuel, but these fractions currently remain in the MOVES database.

For vehicles operating on fuels containing high levels of ethanol, ranging from 70 to 100 percent, the toxic fractions were developed using results of two-day diurnal tests on four 2007 model year flex-fuel vehicles from CRC E-80 program.³⁰ Following typical speciation procedures, the fraction of each compound in a test was first calculated by dividing its emission rates for each compound by the sum of all rates for that test. The percentages for each compound were then averaged across all tests to form the composite profile. The resulting fractions are presented in Table 5-3.

Table 5-1 Data Sources and Estimation Methods Used in Estimation of Toxic Fractions for Evaporative VOCs

Compound	Process	Fraction	Basis for	MOVES Table
		Type	Estimation	
Benzene	Vapor	complex	Adapted from	GeneralFuelRatioExpression for
	venting/refueling		MOBILE6.2	E0 to E20;
	(vapor)			atRatioNonGas for E70-E100
	Fuel leaks/spillage	complex	Adapted from	GeneralFuelRatioExpression for
			MOBILE6.2	E0 to E20;
				atRatioNonGas for E70-E100
2,2,4-	All (except	simple	Speciation	minorHAPratio
trimethylpentane	permeation)		profile	
Ethylbenzene	All (except	simple	Speciation	minorHAPratio
	permeation)		profile	
N-Hexane	All (except	simple	Speciation	minorHAPratio
	permeation)		profile	
Toluene	All (except	simple	Speciation	minorHAPratio
	permeation)		profile	
Xylene	All (except	simple	Speciation	minorHAPratio
	permeation)		profile	
Ethanol	All (except	simple	Speciation	GeneralFuelRatioExpression for
	permeation)	Simple	profile	E0 to E20;
	реппеаноп)		prome	atRatioNonGas for E70-E100
				auxationoligas for E/0-E100

¹For E70 through E100 fuels, the toxic ratios for benzene and ethanol are simple fractions stored in the atRatioNonGas table

Table 5-2 Complex Fractions of VOC for Evaporative Emissions of Benzene Applied to Vehicles Running on Gasoline (E0 to E20)

Pollutant	Process	Equation for Toxic Fraction
D	Vapor venting/Refueling (vapor)	(-0.03420*OXY - 0.080274*RVP + 1.4448)*BNZ/100
Benzene	Fuel Leaks/Spillage	(-0.03420*OXY - 0.080274*RVP + 1.4448)*BNZ/100

Table 5-3 Toxic Fractions for Evaporative VOC Emissions, for Vapor-venting and Refueling-Spillage Processes

Pollutant	Ethanol Level					
	0.0% (E0)	10% (E10)	15% (E15)	20% (E20) ³	70- 100% (E85)	
Ethanol ¹	0.000	0.119	0.194	0.223	0.610	
2,2,4- Trimethylpentane	0.020	0.034	0.053	0.043	0.008	
Ethyl Benzene	0.025	0.017	0.017	0.016	0.001	
N-Hexane	0.022	0.025	0.007	0.019	0.013	
Toluene	0.096	0.143	0.141	0.087	0.016	
Xylene	0.080	0.064	0.057	0.071	0.007	
Benzene	Table 5-2				0.0066	

Notes:

5.1.2 Permeation

The composition of VOCs emitted through permeation differs substantially from that of hydrocarbons emitted through other processes. Work to better characterize these permeation emissions was conducted by Southwest Research Institute for EPA and the Coordinating Research Council in the CRC E-77-2b and E-77-2c test programs. ^{56,57} Data from 3-day diurnal tests on vehicles meeting Tier 1 and near-zero evaporative emission standards were used. Fractions representing emissions of toxic compounds relative to total VOC were estimated for E0, E10 and E20 fuels by averaging data from fuel formulations with varying vapor pressures. To estimate toxic fractions for vehicles operating on fuels containing 15 percent ethanol, the fractions for E10 and E20 fuels were linearly interpolated for ethanol levels of 15 percent. To estimate toxic fractions for vehicles operating on fuels containing 15 percent ethanol, the fractions for E10 and E20 fuels were linearly interpolated for ethanol levels of 15 percent. Toxic fractions are shown in Table 5-4, for all compounds except benzene.

For benzene, the diurnal emissions equation from MOBILE6.2 was used to calculate the permeation fraction $f_{\text{benz,permeation}}$, since it accounts for changes in oxygenate, vapor pressure and fuel benzene levels, as shown in Equation 21.⁵⁸ However, a study of permeation emissions

¹Ethanol toxic fraction is estimated 0.0119×ETOH (The ethanol percent volume in the fuel) using the GeneralFuelRatioTable for E0 to E20 fuels,

²The toxic ratios for Ethanol and Benzene for E70-E100 percent fuel are simple ratios stored in the atRatioNonGas table

³MOVES versions after MOVES2014a do not estimate emissions for vehicles using E20 fuel, but these fractions currently remain in the MOVES database.

suggests that the fraction of benzene from permeation is about 1.77 times higher than the ratio associated with evaporation. ⁵⁹ Thus the diurnal emissions algorithm was multiplied by 1.77.

$$f_{benzene,permeation} = 1.77 \times \left[\frac{(-0.0285 \times OXY(wt\%) - 0.080274 \times RVP + 1.3758) \times benzene}{100} \right] \quad \begin{array}{c} \textbf{Equation} \\ \textbf{21} \end{array}$$

$$= 1.77 \times \left[\frac{(-0.0285 \times (ETOHVolume \times 0.3653) - 0.080274 \times RVP + 1.3758) \times benzene}{100} \right]$$

Where:

Oxy (wt%) = oxygenate fuel content, weight percentage (%)

RVP = reid vapor pressure (psi)

Benzene = benzene fuel content, volume percentage (%)

ETOHVolume = ethanol fuel content, volume percentage (%)

VolToWtPercent = 0.3653 = Parameter to convert ETOHVolume to Oxy (wt%). See Fuel Effects²² report for documentation.

In MOVES, the permeation values are stored in the same location as the toxic values for other evaporative processes. Ethanol and Benzene are stored in the GeneralFuelRatioExpression Table for E0 through E20 fuels, and the atRatioNonGas Table for E70-E100 fuels. The other toxics (2,2,4-Trimethylpentane, Ethylbenzene, Hexane, Toluene and Xylene) are stored in the minorHAPratio table.

Table 5-4 Toxic Fractions Representing Permeation Emissions as Components of Total VOC Emissions, by Ethanol Level (Source: CRC E-77-2b and CRC E-77-2c)

Pollutant	Ethanol Level					
	0.0% (E0)	10% (E10)	15% (E15)	20% (E20) ²	70-100% (E85)	
Ethanol	0.000	0.202	0.2694	0.3296	0. 6104 ¹	
2,2,4-Trimethylpentane	0.036	0.024	0.0172	0.0107	0. 00831	
Ethylbenzene	0.003	0.001	0.0017	0.0019	0. 0012 ¹	
Hexane	0.050	0.065	0.0472	0.0308	0. 01281	
Toluene	0.110	0.101	0.0666	0.0354	0. 0161 ¹	
Xylene(s)	0.016	0.011	0.0127	0.0140	0. 00731	
Benzene	Equation 21				0. 00661	

Note:

For ethanol levels of 70-100 percent, no permeation data were available. Thus, the toxic fraction for non-permeation evaporative emissions was also applied to permeation.

¹ Identical to fractions for the vapor-venting process, based on CRC E-80 program (Table 5-3).

² MOVES versions after MOVES2014a do not estimate emissions for vehicles using E20 fuel, but these fractions currently remain in the MOVES database.

5.2 Diesel Vehicles

For diesel-fueled vehicles, evaporative emissions are estimated for the refueling-spillage process only. In MOVES3, we updated the diesel spillage air toxic fractions to be based on the liquid diesel fuel (Profile 95120 Liquid Diesel - California composite) stored in the SPECIATE database. His profile is also used to speciate the other VOC emissions from diesel refueling spillage. The fractions are shown in Table 5-5 and are stored in the minorHAPratio table, except for benzene which is stored in the atRatioNonGas table.

Table 5-5 Toxic Fractions for the Fuel-Spillage Process for Diesel Fuel

Pollutant	Toxic fraction		
2,2,4-Trimethylpentane	0		
Ethyl Benzene	0.00103		
N-Hexane	0		
Toluene	0.00235		
Xylene	0.00706		
Benzene	0		
Naphthalene gas	0.00048		

6 Crankcase Emissions

Crankcase emissions are modeled as a ratio of the exhaust emissions. Discussion of the ratios used to estimate THC, CO, NOx, and PM crankcase emissions can be found in the light-duty⁶⁰ and heavy-duty⁶¹ emission rate reports. MOVES calculates gaseous crankcase toxics as a fraction of the exhaust toxics described in the sections above. Particulate toxics are handled differently. MOVES does not model crankcase dioxins, furans, or many metals. The details on crankcase emissions are discussed in the following sections.

6.1 Volatile Organic Compounds

Table 1-1 lists the VOC toxics modeled in MOVES which are also modeled from crankcase emission processes. MOVES models the crankcase emissions of these toxics by multiplying the exhaust emissions of these species by the non-methane hydrocarbon (NMHC) crankcase emission fraction listed in the light-duty and heavy-duty emissions reports. For example, the NMHC crankcase/exhaust fraction for light-duty gasoline (1969 and later model year) is 0.013. Thus, crankcase emissions for 1,3-butadiene are calculated as 1.3 percent of the exhaust emissions of 1,3-butadiene. Similar calculations are applied to all VOC toxic emissions. The crankcase emission ratios are stored in the MOVES table *CrankcaseEmissionRatio*, which differentiates the factors according to pollutant, process, model year range, source type and fuel type.

6.2 Polycyclic Aromatic Hydrocarbons

Crankcase gaseous PAHs are modeled in a similar fashion as the crankcase VOC toxic emissions. The PAH crankcase emissions are modeled as a fraction of the tailpipe exhaust gaseous PAH emissions, with factors stored in the crankcaseEmissionRatio table. The PAH crankcase emission factors are the same as the NMHC crankcase emission factors (e.g. 0.013 for 1969 and later gasoline vehicles, and 0.39 for 2010 and later heavy-duty diesel vehicles).

Because the OC/PM speciation can be substantially different between crankcase emissions and exhaust emissions, we use a different approach for these calculations. To estimate crankcase particulate PAH emissions, MOVES applies the same PAH/OC fractions developed for exhaust emissions to the crankcase OC emissions. The PAH/OC ratios are stored in the pahParticleRatio table for the crankcase emission processes (15, 16, and 17). For example, because conventional diesel crankcase emissions have a higher OC/PM composition than the tailpipe exhaust emissions, MOVES models elevated particulate PAH emissions in crankcase PM_{2.5} emissions compared to tailpipe PAH emissions. Research on conventional diesel vehicles validates that PM emissions from the crankcase are more enriched with PAHs than emissions from the exhaust. 62

6.3 Metal and Dioxin Emissions

MOVES models crankcase metal emissions for the metal species included in the PM_{2.5} exhaust speciation profiles, such as iron and aluminum, which are also applied to crankcase emissions. Details on speciation of crankcase emissions are included in the speciation report.²¹ MOVES

does not produce crankcase emission rates for metals that are not included in the speciation profiles such as arsenic, mercury and other metals listed in Table 1-4. Similarly, MOVES does not estimate dioxin and furan emissions from crankcase emissions, assuming that the emissions from crankcase are negligible compared to exhaust emissions.

Appendix A. Development of Motor Vehicle Emission Factors for Chromium

In 2010, the EPA's National Vehicle and Fuel Emissions Laboratory (NVFEL) collected particulate matter (PM) and volatile organic compound (VOC) exhaust samples, as well as CO, NO_x, CO₂, and CH₄ samples from a 2008 3.5L V6 Chevrolet Impala flex fuel light-duty gasoline vehicle. This testing also included direct Cr(VI) measurements.

The Impala had a beginning odometer reading of 38,934 miles and was tested using E10 gasoline. The vehicle test procedure used four sample bags and the LA92 "unified" dynamometer driving schedule. 63 The bags in this study represent the following conditions:

Bag 1 – concentrated cold start compared to FTP (Federal Test Procedure); short distance, low speeds.

Bag 2 – hot and running; longer distance and higher speeds than FTP (represents realistic real world driving).

Bag 3 – hot start; short distance, low speeds.

Bag 4 – hot and running; long distance.

PM was collected on four (labeled A-D) pre-cleaned and prepared filter media per bag. The PM filter samples labeled D were sent to the Wisconsin State Laboratory of Hygiene at the University of Wisconsin-Madison for chromium metal speciation. Total and hexavalent chromium was measured in extracts of filter-collected PM sent from NVFEL. Detection limits were in the <0.2 ng/filter range. A comparison of 47mm filter collection substrates was performed using Polyvinyl Chloride (PVC) and bicarbonate-impregnated Mixed Cellulose Ester (MCE) filters. Total chromium was analyzed by SF-ICPMS (Sector Field Inductively Coupled Plasma Mass Spectrometry) and Cr(VI)was analyzed by Inductively Coupled (IC)-post-column derivation. The Cr(VI) results obtained using PVC collection substrates were below the detection limit, with the exception of the tunnel blanks, and thus not listed in this memo. The extractable total chromium levels in the filters and bicarbonate were at such a level that swamp any signal from the PM, making the ICPMS data useless. However, the Cr(VI) data from the MCE filters analyzed by IC could be used to develop new emission rates as described below.

Spike and blank studies were performed. Spike studies had a recovery between 93-104 percent, indicating the matrix did not interfere with the chromium results. The Cr(VI) MCE filter results were blank corrected by subtracting the mean background value of 0.298 ng/filter (standard deviation±0.098 ng/filter; 95 percent confidence interval±0.157). The 95 percent confidence interval was calculated from student's *t*-distribution as a function of the probability and degrees of freedom and multiplied by the standard deviation over the square root of the number of blanks.

Cr(VI) speciation results and emission rates are reported in Table A-1 along with the corresponding distance driven per sample. The emission rates were calculated by dividing the blank-corrected Cr(VI) MCE mass/filter by the distance driven per sample and multiplying by a factor representing the CVS (constant volume sampler) volume over the individual filter sample

volume (NVFEL filter sample D was used for each bag). This factor was used because all exhaust was not passed through the collection filter during the test.

$$Emission \ Rate = \frac{blank corrected \ Cr(VI)MCE \ mass/filter}{distance} \times \frac{CVS \ volume}{Sample \ volume}$$

The overall emission rate in Table A-1 is a composite average of the total Cr(VI) measured divided by the total distance of the test and then multiplied by the sum of CVS volumes/sum of filter sample volumes.

Sample/bag number	Cr(VI) (ng/filter)	Mean IC Blank± Std Deviation (ng/filter)	Blank- corrected Cr(VI) (ng/filter)	CVS Volume (scf at 68°F)	Sample Volume (scf at 68°F)	Distance (miles)	Emission Rate (g/mile)
1	0.792	0.298±0.098	0.49	1666.87	7.675	1.194	8.9x10 ⁻⁸
2	0.493	0.298 ± 0.098	0.20	6280.73	28.815	8.612	5.1x10 ⁻⁹
3	0.488	0.298 ± 0.098	0.19	1682.74	7.711	1.186	3.5x10 ⁻⁸
4	0.508	0.298±0.098	0.21	6281.82	28.894	8.620	5.3x10 ⁻⁹
Overall			1 1	15012.2	73.10	10.61	1.2×10-8

Table A-1 Cr(VI) Emission Rates from an Onroad Gasoline Engine

Direct Cr(VI) emission factors were not measured from a diesel engine. To develop on-road diesel emission factors, the total chromium from diesel engines was multiplied by the ratio of Cr(VI) to total chromium from onroad gasoline emissions (0.29%). The Cr(VI) onroad emission rates are shown in Table A-1. The total chromium emission factor for gasoline comes from the Kansas City Particulate Matter Characterization Study (4.07x10⁻⁶ g/mi).³² The KCVES program sampled 99 vehicles for chemical composition from which a total chromium emission factor of 4.07x10⁻⁶ grams/mile was developed.³⁵ This average grams/mile rate was calculated by averaging the metal measured in Bag 2 of the LA92 driving schedule test with a weighted-average computed using vehicle miles traveled (VMT).

Diesel Cr(VI) emission factors are calculated for diesel engines based on total chromium from test programs conducted on diesel engines before³⁹ and after implementation of EPA's 2007⁴², and 2010⁴³ heavy-duty highway rule which reduced PM emissions from heavy-duty diesel vehicles, as outlined in the following equations

Cr(VI) Pre-2007 On-road Diesel Emission Factor

$$EF = Total\ Cr\ EF_{Pre2007\ diesel} \times \frac{Gasoline\ Cr(VI)EF}{Total\ Cr\ EF_{gasoline}} = 6.8x10^{-6} \frac{g}{mi} \times \frac{1.2x10^{-8} \frac{g}{mi}}{4.07x10^{-6} \frac{g}{mi}} = 6.8x10^{-6} \frac{g}{mi} \times 0.29\%$$

$$= 2.0x10^{-8} \frac{g}{mi}$$

Cr(VI) 2007 to 2009 On-road Diesel Emission Factor

$$EF = Total\ Cr\ EF_{2007-2009\ diesel} \times \frac{Gasoline\ Cr(VI)EF}{Total\ Cr\ EF_{gasoline}} = 2.01x10^{-6} \frac{g}{mi} \times 0.29\% = 5.9x10^{-9} \frac{g}{mi}$$

Cr(VI) 2010 and Later On-road Diesel Emission Factor

$$EF = Total\ Cr\ EF_{2010+diesel} \times \frac{Gasoline\ Cr(VI)EF}{Total\ Cr\ EF_{gasoline}} = 7.3x10^{-7} \frac{g}{mi} \times 0.29\% = 2.2x10^{-9} \frac{g}{mi}$$

A Cr(VI) emission factor for vehicles using compressed natural gas is calculated by multiplying the overall Cr(VI) emission factor from Table A-1 by the ratio of total chromium from CNG transit buses⁶⁴ verses gasoline light-duty vehicle engines (from KCVES study).

Cr(VI) Compressed Natural Gas (CNG) Emission Factor

$$EF = Total\ Cr\ EF_{CNG} \times \frac{Gasoline\ Cr(VI)EF}{Total\ Cr\ EF_{agsoline}} = 7.0x10^{-8} \frac{g}{mi} \times 0.29\% = 2.1x10^{-10} \frac{g}{mi}$$

A summary of the results for Cr(VI) emission factors is presented in Table A-2. While these results are based on measured Cr(VI), the results are limited by the following:

- Emissions from only one vehicle were measured, so the data do not provide information regarding variability among vehicles.
- No Cr(VI) measurements have been made for diesel and CNG vehicles or engines.

Table A-2 Summary: Cr(VI) Emission Factors

	Emission Factor	Units
Onroad gasoline (MY2008)	$1.2x10^{-8}$	grams/mile
Onroad diesel (pre-2007) Onroad diesel (2007-2009 and	$2.0x10^{-8}$	grams/mile
later)	5.9×10^{-9}	grams/mile
Onroad diesel (2010 and later)	$2.2x10^{-9}$	grams/mile
CNG heavy-duty vehicles	2.1×10^{-10}	grams/mile

Appendix B. Development of Motor Vehicle Emission Factors for Mercury

Calculation of Mercury Emission Factors from Vehicle Tests

In 2005, the USEPA National Exposure Research Laboratory (NERL) collected mercury (Hg) samples in the raw exhaust from 14 light-duty gasoline vehicles and two heavy-duty diesel vehicles. The work plan for this project includes details of the methods used that are not reproduced here including quality assurance and quality control for Hg collection and analysis. This information can be obtained from EPA upon request. Briefly, mercury and regulated pollutant data were collected during two sets of three consecutive LA92 drive cycles for each vehicle. The morning set of LA92 cycles began with one 'cold start' and the afternoon set of three LA92 cycles began with a 'hot start'. The intake air was filtered through charcoal to greatly reduce background mercury concentrations entering the vehicle intake. Separate sample lines were used for gaseous and particulate mercury species. Samples analyzed for mercury were drawn from raw exhaust at a constant flow rate and fixed dilution. Carbon dioxide measurements were also taken in the exhaust stream where mercury samples were collected.

Mercury samples were collected in the raw exhaust since previous data suggested that mercury levels might be sufficiently low to challenge mercury detection limits. This sampling method imposed a challenge in calculating emission factors since it assumes that the exhaust flow rate from the vehicle is constant. Calculation of exhaust flow and its application to the development of mercury emission rates is described below.

Evaporative losses of mercury from motor vehicles and loss of mercury during refueling were not measured. The emission of mercury through evaporative processes is expected to be negligible compared with that expected from exhaust emissions.

A description of the vehicles tested for which data were used in developing emission rates is provided in Table B-1. The data collected from these vehicles in diluted exhaust in the constant volume sampler (CVS) included THC, carbon dioxide (CO₂), nitrogen oxides (NO_x), methane (CH₄), and carbon monoxide (CO). In raw, undiluted exhaust, data collected included elemental and total gas-phase mercury, particulate mercury and CO₂. Gas-phase mercury was also measured in the intake air. Total air flow was measured for all sampling systems and corrected to standard temperature and pressure conditions. The data streams had different reporting frequencies, all due to the nature of the instrumentation. The dilute measurement of the standard emission gases (THC, CO₂, NO_x, CH4, and CO), CVS flows, and vehicle speed were reported at 1 Hertz. The gas-phase mercury samples were analyzed at 2.5-minute intervals and particle-phase mercury samples were collected cumulatively for the duration of three consecutive LA92 cycles. Gas-phase elemental mercury in the engine intake air was measured at five-minute intervals.

Table B-1 Vehicles Tested for Mercury Emissions

Model Year	Make	Model	Fuel Type	Odometer	Cylinders	Displacement
				(mi)	-	(L)
2005	MERCURY	GRAND MARQUIS LS	Gasoline	9,953	8	4.6
2005	FORD	MUSTANG CONVERTIBLE	Gasoline	5,424	6	4.0
2003	SATURN	L 200	Gasoline	29,667	4	2.2
2002	HONDA	ACCORD EX	Gasoline	51,824	4	2.3
2001	HONDA	ACCORD EX	Gasoline	88,611	4	2.3
2001	CHRYSLER	PT CRUISER	Gasoline	54,010	4	2.4
2000	CHEVROLET	SUBURBAN	Gasoline	39,787	8	6.0
2000	JEEP	CHEROKEE SPORT	Gasoline	48,468	6	4.0
1999	FORD	F250 XLT	Diesel	113,897	8	7.3
1999	FORD	F250 XLT SD	Diesel	109,429	8	7.3
1998	HONDA	CIVIC DX	Gasoline	204,983	4	1.6
1994	CHEVROLET	SILVERADO	Gasoline	129,521	8	5.7
1992	CHEVROLET	S10 BLAZER	Gasoline	162,249	6	4.3
1991	HONDA	ACCORD EX	Gasoline	143,289	4	2.2
1987	CHRYSLER	FIFTH AVENUE	Gasoline	72,573	8	5.2
1984	FORD	F150 PICKUP	Gasoline	36,727	8	5.8

Exhaust flow was integrated at the same reporting frequency as the mercury exhaust values for a particular test and then used to calculate total, elemental, and reactive gas-phase mercury mass emissions. The intake air mercury values were typically collected at half the frequency of the mercury exhaust values and used to correct exhaust measured values that are reported at higher frequencies. The particulate matter measurements were filter-based, test-level measurements and were corrected in that manner.

Calculation of Emission Rates

Emission rates were calculated separately for elemental gas-phase mercury, reactive gas-phase mercury and particulate mercury. Elemental gas-phase mercury in the exhaust was corrected for the intake air concentration of elemental mercury. To estimate the gas-phase mercury concentration in dilute exhaust from the measured mercury in raw exhaust, the dilution factor was applied. For light-duty gasoline vehicles, the dilution factor equation found in 40 CFR 90.426 (d) was used:

Dilution factor =
$$13.4 / ([CO_2\%] + ([THC, ppm] + [CO, ppm])* 0.0001)$$

Exhaust flow = (CVS flow / dilution factor)

Exhaust flow calculation was initiated when the analytical equipment indicated that the dilute exhaust CO₂ concentration was greater than the background CO₂ concentration.

To calculate exhaust flow for the diesel vehicles, the dilution factor was calculated by simply dividing CO₂ in the raw exhaust by CO₂ in the CVS. This method was used because diesel engines operate across a very wide range of fuel to air mixtures and the CFR method described above was not appropriate.

Determination of Reactive Gas Mercury Mass in Exhaust

Reactive gas-phase mercury (RGM) was calculated by subtracting elemental gas-phase mercury measurements from total gas-phase mercury measurements. RGM values were typically small and therefore influenced by the variability in the elemental mercury measurements. Negative RGM values for a given measurement period were observed. Values for which there was not a positive RGM measurement were treated as non-detects and were nulled in the aggregation of RGM values for the test. The measurement uncertainty for gas-phase elemental mercury was estimated from quantitative recovery of injections of known amounts of mercury into the sampling system. The uncertainty in measuring elemental mercury was applied to the total gas-phase and elemental gas-phase measurements to determine when the RGM value was above the measurement uncertainty. Values within the measurement uncertainty were not included in the emission factor calculation.

Calculating Weighted Emission Test Results

Highway vehicles were tested on the LA92 cycle; a more aggressive chassis-dynamometer test similar in concept to the Federal Test Procedure's (FTP) UDDS or LA4. Like the FTP, the LA92 includes a cold start, a hot start, and a hot stabilized phase using identical drive schedules for the starts. We considered it appropriate to calculate a weighted emission factor (representing cold start and hot start driving) for each vehicle in the same manner as the FTP, using the equation below for each test (a test consisting of all six LA92 cycles performed on each vehicle).

We summed the gas-phase mercury mass emissions for the first phase (300 seconds) of the morning test and last phase (1,135 seconds) of the individual LA92 drive schedules for all the tests (e.g., 'hot stabilized emissions'), divided by the total distance covered in these phases and multiplied by 0.43. We also summed the sum of the mass gas-phase mercury emissions of the first phase of the afternoon test and last phase (1,135 seconds) of all the tests, divided by the total distance covered in these phases and multiplied by 0.57. The two terms were summed to calculate a test level emission rate for each of the gasoline powered vehicles.

The equation used to calculate test-level emission rates is as follows:

$$\bar{E}_{\text{Hg}} = 0.43 \left(\frac{C+R}{C_m + R_m} \right) + 0.57 \left(\frac{H+R}{H_m + R_m} \right)$$

Where:

 $E_{\rm Hg}$ = mean aggregate emission rate (g/mi),

C = mercury mass collected in the first 300 seconds of the first morning test ('cold start', g), C_m = distance covered in the cold start phase (mi),

R = mercury mass collected in the last 1,135 seconds of all six cycles of the LA92 ('hot stabilized', g),

 R_m = cumulative distance covered in all six cycles of the LA92 ('hot stabilized', mi),

H = mercury mass collected in the first 300 seconds of the first afternoon test ('hot start', g),

 H_m = distance covered by the hot start (mi).

It should be noted that the 'hot start' in the afternoon typically occurred after the vehicle had been off for at least 1 hour, making this start closer to a 'cold start' than 'hot start'. Since the true cold start emissions were slightly higher than hot start emissions, it is expected that this approach would bias the emission factors high by a small amount, relative to the value expected for a cycle composite.

Particulate mercury emissions could not be apportioned into modes of operation in similar manner because filters were collected across all three LA92 cycles and could not be parsed into the three phases. A test-level composite emission rate was calculated by multiplying the morning particulate mercury emission rate by 0.43 and the afternoon particulate mercury emission rate by 0.57 and adding the two values together.

The average of emission factors across vehicles was calculated for each form of mercury and is reported in Table B-2. A simple average was used since the data did not suggest that mercury concentrations varied by vehicle age, mileage, displacement or other factors.

Mercury emission factors for on-road diesel engines were obtained from the first 715 seconds of the morning and afternoon tests on the Ford F250 XLT SD; data from the second diesel vehicle could not be used. The first 715 seconds is approximately half of the first of the three LA92 drive cycles that made up a single test. The truncation of the test was due to sample flow problems in the mercury sampling manifold due to particulate matter restricting flow across the particulate matter filters. Graphical analysis of exhaust flow indicated that they appeared nominal during the first LA92 cycle. We decided that only using measurements collected before 715 seconds in both tests provided the most reliable data.

Nonroad grams per gallon emission factors in Table B-2 were calculated from the onroad factors using a fuel economy estimate of 17 miles per gallon for the gasoline vehicle and 19 for the diesel vehicle.

Table B-2 Mercury Emission Factors from Mobile Sources

Table B-2 Mercury Emission Factors from Mobile Sources					
Source Category	Pollutant	Pollutant	Emission	Units	
		ID	Rate		
Gasoline motor vehicles	Elemental gas- phase	200	1.1E-07	grams/mile	
	Reactive gas-phase	201	9.9E-09	grams/mile	
	Particulate phase	202	4.0E-10	grams/mile	
Diesel motor vehicles	Elemental gas- phase	200	6.2E-09	grams/mile	
	Reactive gas-phase	201	3.2E-09	grams/mile	
	Particulate phase	202	1.6E-09	grams/mile	
Gasoline nonroad engines	Elemental gas- phase	200	1.8E-06	grams/gallon	
	Reactive gas-phase	201	1.7E-07	grams/gallon	
	Particulate phase	202	6.9E-09	grams/gallon	
Diesel nonroad engines	Elemental gas- phase	200	1.2E-07	grams/gallon	
	Reactive gas-phase	201	6.2E-08	grams/gallon	
	Particulate mercury	202	3.2E-08	grams/gallon	

7 References

¹ Code of Federal Regulations, 40: Chapter 1, Subchapter C, Part 51, Subpart F, 51100, 1986.

² Chemical Abstracts Service, A division of the American Chemical Society. http://www.cas.org.

³ USEPA (2016). *Air Toxic Emissions from On-road Vehicles in MOVES2014*. EPA-420-R-16-016. Office of Transportation and Air Quality. US Environmental Protection Agency. Ann Arbor, MI. November 2016. http://www.epa.gov/moves/moves-technical-reports.htm.

⁴ USEPA (2020). Fuel Supply Defaults: Regional Fuels and the Fuel Wizard in MOVES3. EPA-420-R-20-017. Office of Transportation and Air Quality. US Environmental Protection Agency. Ann Arbor, MI. November 2020. https://www.epa.gov/moves/moves-technical-reports.

⁵ USEPA (2017). Speciation and Toxic Emissions from Onroad Vehicles, and Particulate Matter Emissions from Light-Duty Gasoline Vehicles in MOVES201X - Draft Report. Draft report and peer-review documents. Record ID 328810. EPA Science Inventory. September 2017. https://cfpub.epa.gov/si/si public record report.cfm?dirEntryId=328810.

⁶ USEPA (2000). "Control of Air Pollution From New Motor Vehicles: Tier 2 Motor Vehicle Emissions Standards and Gasoline Sulfur Control Requirements", 65 FR 6698, February 10, 2000.

⁷ USEPA (2014). Control of Air Pollution from Motor Vehicles: Tier 3 Motor Vehicle Emission and Fuel Standards; Final Rule. *Federal Register*. Vol. 79, No. 81. April 28, 2014. Page 23417.

⁸ USEPA. (1993). Final Regulatory Impact Analysis for Reformulated Gasoline. December 13, 1993. Available at https://nepis.epa.gov/Exe/ZyPURL.cgi?Dockey=P1002607.txt

⁹ Sierra Research (2010). *Development of Emission Rates for the MOVES Model*. Prepared for the U. S. Environmental Protection Agency by Sierra Research, Sacramento, CA, July 2, 2010. Sierra Research Report No. SR2010-07-01.

¹⁰ Southwest Research Institute (2007). Flex Fuel Vehicles (FFVs): VOC/PM Cold Temperature Characterization When Operating on Ethanol (E10, E70, E85). Prepared for U. S. Environmental Protection Agency.

¹¹ Environment Canada (2007). Comparison of Emissions from Conventional and Flexible Fuel Vehicles Operating on Gasoline and E85 Fuels. ERM Report No. 05-039. Emissions Research Division.

¹² Durbin T. D., Miller J. W., Younglove T., Huai T., Cocker K. (2007). Effects of fuel ethanol content and volatility on regulated and unregulated exhaust emissions for the latest technology gasoline vehicles. *Environmental Science and Technology* 41, 4059-4064.

¹³ USEPA (1995). MOVES Toxic Inputs 5.xls. Office of Research and Development, unpublished data.

¹⁴ USEPA (2011). *EPA Announces E15 Partial Waiver Decision* EPA-420-F-11-003. Office of Transportation and Air Quality. US Environmental Protection Agency. January 2011. https://nepis.epa.gov/Exe/ZyPDF.cgi?Dockey=P1009LS8.pdf.

¹⁵ USEPA (2013). EPAct/V2/E-89: Assessing the Effect of Five Gasoline Properties on Exhaust Emissions from Light-Duty Vehicles certified to Tier-2 Standards: Final Report on Program Design and Data Collection. EPA-420-R-13-004. Assessment and Standards Division, office of Transportation and Air Quality, Ann Arbor, MI; National Renewable Energy Laboratory, Golden, CO; Coordinating Research Council, Alpharetta, GA. April, 2013.

¹⁶ USEPA (2013). Assessing the Effect of Five Gasoline Properties on Exhaust Emissions from Light-Duty Vehicles certified to Tier-2 Standards: Analysis of Data from EPAct Phase 3 (EPAct/V2/E-89). Final Report. EPA-420-R-13-002. Assessment and Standards Division, Office of Transportation and Air Quality, Ann Arbor, MI. April, 2013.

- ¹⁷ Neter, J.: Kutner, M.H.; Nachtsheim, C.J.; Wasserman, W. (1996). *Applied Linear Statistical Models*. Fourth Ed., Irwin, Chicago. (Section 7.5).
- ¹⁸ West, Brady T.; Welch, Kathleen B.; Galecki, Andrzej T. (2007). *Linear Mixed Models: A Practical Guide Using Statistical Software*. Taylor and Francis CRC Press. 376 pp.
- ¹⁹ SAS Institute Inc. (1990). SAS/STAT User's Guide, Version 6, Fourth Ed., Vol. 2. Chapter 25, The LIFEREG Procedure. Pp. 997-1026. Cary, NC.
- ²⁰ Neter, J.; Kutner, M.H.; Nachtsheim, C.J.; Wasserman, W. (1996). *Applied Linear Statistical Models*. Fourth Ed., Irwin, Chicago.
- ²¹ USEPA (2020). *Speciation of Total Organic Gas and Particulate Matter Emissions from Onroad Vehicles in MOVES3*. EPA-420-R-20-021. Office of Transportation and Air Quality. US Environmental Protection Agency. Ann Arbor, MI. November 2020. https://www.epa.gov/moves/moves-technical-reports.
- ²² USEPA (2020). Fuel Effects on Exhaust Emissions from Onroad Vehicles in MOVES3. EPA-420-R-20-016. Office of Transportation and Air Quality. US Environmental Protection Agency. Ann Arbor, MI. November 2020. https://www.epa.gov/moves/moves-technical-reports.
- ²³ Zarnoch, S.J., and W.A. Bechtold (2000). Estimating mapped-plot forest attributes with ratios of means. *Canadian Journal of Forest Research*. 30: 688-697.
- ²⁴ USEPA (2007). Final Regulatory Impact Analysis: Control of Hazardous Air Pollutants from Mobile Sources. EPA420-R-07-002; U.S. Environmental Protection Agency; Office of Transportation and Air Quality: Ann Arbor, MI. Chapter 6.11. http://www.epa.gov/otaq/toxics.htm.
- ²⁵ USEPA (2011). Spreadsheet "Benz Buta Adjustments.xlsx." EPA Docket EPA-HQ-OAR-2011-0135.
- ²⁶ USEPA (2011). Spreadsheets "Olefins versus Butadiene.xls" and "Benz Buta Adjustments.xls." EPA Docket EPA-HQ-OAR-2011-0135.
- ²⁷ USEPA (2009). Exhaust Emission Profiles for EPA SPECIATE Database: Energy Policy Act (EPAct) Low-Level Ethanol Fuel Blends and Tier 2 Light-Duty Vehicles. EPA-420-R-09-002. Office of Transportation and Air Quality, Assessment and Standards Division, Ann Arbor, MI.
- ²⁸ USEPA (2012). Data Collected in EPAct Fuel Effects Study Pilot Phases. Memorandum from Aron Butler to Docket EPA-HQ-OAR-2011-0135.
- ²⁹ Yanowitz, J., Knoll, K., Kemper, J., Luecke, J., and McCormick R. (2013) *Impact of Adaptation on Flex-Fuel Vehicle Emissions When Fueled with E40*. Environmental Science & Technology, 2013, 47(6), 2990-2997.
- ³⁰ Haskew, H.M., and Liberty, T.F. (2011) *Exhaust and Evaporative Emissions Testing of Flexible-Fuel Vehicles. Final Report.* Coordinating Research Council. CRC Report No. E-80, August, 2011. Available at www.crcao.org.
- ³¹ Long, T., Herrington, J., Hays, M., Baldauf, R., and Snow, R. (2011) *Air Toxic Emission from Passenger Cars Operating on Ethanol Blend Gasoline*. 104th Air and Waste Management Association Annual Conference and Exhibition, 2011, #118, 3053-3058.

- ³³ Fujita, E. M.; Campbell, D. E.; Zielinska, B.; Chow, J. C. (2006). *Kansas City Gasoline Vehicle Characterization Study Effect of Sampling Temperature on Chemical Speciation*. NREL Subcontract Number ACI-5-55500-01. NREL Project ES04-2. May 31, 2006.
- ³⁴ Hays, M. D., et al. (2013). Carbonaceous Aerosols Emitted from Light-Duty Vehicles Operating on Gasoline and Ethanol Fuel Blends. *Environ Sci Technol*, 47 (24), 14502-14509. DOI: Doi 10.1021/Es403096v.
- ³⁵ Sonntag, D. B., R. W. Baldauf, C. A. Yanca and C. R. Fulper (2013). Particulate matter speciation profiles for light-duty gasoline vehicles in the United States. *Journal of the Air & Waste Management Association*, 64 (5), 529-545. DOI: 10.1080/10962247.2013.870096.
- ³⁶ Schauer, J. J., Lough, G. C., Shafer M. M., Christensen W. F., Arndt, M. F., DeMinter, J.T., Park, J-S. (2006). *Characterization of Metals Emitted from Motor Vehicles*. Health Effects Institute Research Report Number 133. (http://pubs.healtheffects.org).
- ³⁷ Gertler, A. W., J. C. Sagebiel, W. A. Dippel and R. J. Farina. (1998). Measurement of dioxin and furan emission factors from heavy-duty diesel vehicles. *J. Air and Waste Manage. Assoc.* 48: 276-278.
- ³⁸ USEPA (2000). *Draft Final Assessment: Exposure and Human Health Reassessment of 2,3,7,8-Tetrachlorodibenzo-p-Dioxin (TCDD) and Related Compounds. Volume 2: Sources of Dioxin-like Compounds in the United States*. Office of Research and Development, National Center for Environmental Assessment, Washington, D.C. Report No. EPA/600/P-00/001Bb, September 2000.
- ³⁹ Hsu ,Y., and Mullen, M. (2007). *Compilation of Diesel Emissions Speciation Data*. Prepared by E. H. Pechan and Associates for the Coordinating Research Council. CRC Contract No. E-75, October, 2007. Available at www.crcao.org.
- ⁴⁰ USEPA (2011). Final Rulemaking: Greenhouse Gas Emissions Standards and Fuel Efficiency Standards for Medium- and Heavy-Duty Engines and Vehicles. Final Regulatory Impact Analysis, Chapter 5 (RIA) (PDF) (553 pp, EPA-420-R-11-901, August 2011)
- ⁴¹ USEPA (2016). Greenhouse Gas Emissions and Fuel Efficiency Standards for Medium- and Heavy-Duty Engines and Vehicles Phase 2. Federal Register 81(206): 73478-74274. Available at https://www.gpo.gov/fdsys/pkg/FR-2016-10-25/pdf/2016-21203.pdf
- ⁴²Khalek, I., Bougher, T., and Merritt, P. M. (2009). Phase 1 of the Advanced Collaborative Emissions Study. Prepared by Southwest Research Institute for the Coordinating Research Council and the Health Effects Institute, June 2009. Available at www.crcao.org.
- ⁴³ Khalek, I., Blanks, M., and Merritt, P. M. (2013). Phase 2 of the Advanced Collaborative Emissions Study. Prepared by Southwest Research Institute for the Coordinating Research Council and the Health Effects Institute, November 2013. Available at www.crcao.org.
- ⁴⁴ USEPA (2020). SPECIATE5.1. https://www.epa.gov/air-emissions-modeling/speciate
- ⁴⁵Sierra Research, 2010. Development of Emission Rates for the MOVES Model. Prepared for the U. S. Environmental Protection Agency by Sierra Research, Sacramento, CA, July 2, 2010. Sierra Research Report No. SR2010-07-01.

³² USEPA (2008). *Kansas City PM Characterization Study: Final Report*. EPA420-R-08-009. Office of Transportation and Air Quality. Assessment and Standards Division . http://www.epa.gov/oms/emission-factors-research/420r08009.pdf.

- ⁴⁷ Liu, Z. G., J. C. Wall, N. Ottinger and D. McGuffin (2015). Mitigation of PAH and Nitro-PAH Emissions from Nonroad Diesel Engines. *Environ Sci Technol*. DOI: 10.1021/es505434r.
- ⁴⁸ Laroo, A. Christopher, Charles R. Schenk, L. James Sanchez, Joseph McDonald, Peter L. Smith (2012). Emissions of PCDD/Fs, PCBs, and PAHs from legacy on-road heavy-duty diesel engines. *Chemosphere* 89: 1287 1294.
- ⁴⁹ Laroo, A. Christopher, Charles R. Schenk, L. James Sanchez, and Joseph McDonald. (2011). Emissions of PCDD/Fs, PCBs, and PAHs from a Modern Diesel Engine Equpped with Catalyzed Emission Control Systems. Environmental Science & Technology 45: 6420 6428.
- ⁵⁰ Ayala, A., Gebel, M., Okamoto, R., Rieger, P. et al. (2003). "Oxidation Catalyst Effect on CNG Transit Bus Emissions," Society of Automotive Engineers, SAE Technical Paper 2003-01-1900, 2003. (http://papers.sae.org/2003-01-1900).
- ⁵¹ Kado, N. Y.; Okamoto, R. A.; Kuzmicky,, P. A.; Kobayashi, R.; Ayala, A.; Gebel, M. E.; Rieger, P. L.; Maddox, C.; Zafonte, L.; (2005). Emissions of Toxic Pollutants from Compressed Natural Gas and Low Sulfur Diesel-Fueled Heavy-Duty Transit Buses Tested over Multiple Driving Cycles. Environmental Science & Technology 2005 39 (19), 7638-7649
- ⁵² Okamoto, R. A.; Kado, N. Y.; Ayala, A.; Gebel, M.; Rieger, P.; Kuzmicky, P. A.; Kobayashi, R.; (2004) Chemical and Bioassay Analyses of Emissions from Two CNG Buses with Oxidation Catalyst. http://www.arb.ca.gov/research/veh-emissions/cng-diesel/cng-diesel.htm.
- ⁵³ USEPA (2020). Evaporative Emissions from Onroad Vehicles in MOVES3. EPA-420-R-20-012. Office of Transportation and Air Quality. US Environmental Protection Agency. Ann Arbor, MI. November 2020. https://www.epa.gov/moves/moves-technical-reports.
- ⁵⁴ USEPA (2002). Technical Description of the Toxics Module for MOBILE6.2 and Guidance on Its Use for Emission Inventory Preparation. EPA420-R-02-011. U.S. Environmental Protection Agency; Office of Transportation and Air Quality: Ann Arbor, MI. Available at http://www.epa.gov/otaq/m6.htm.
- ⁵⁵C.E. Lindhjem (2008). Emission Profiles for EPA SPECIATE Database, EPA Contract No. EP-C-06-094, Work Assignment No. 1-7, ENVIRON International Corporation, January 31, 2008.
- ⁵⁶Southwest Research Institute (2009). Evaporative Emissions Breakdown Including Permeation Effects and Diurnal Emissions on Aging Enhanced Evaporative Emissions Certified Vehicles (CRC E-77-2b). Prepared by Harold M. Haskew and Thomas F. Liberty, Harold Haskew and Associates, Inc.; submitted to U. S. EPA, December, 2009. Available at www.crcao.org.
- ⁵⁷ Coordinating Research Council (2010). Study to Determine Evaporative Emission Breakdown, Including Permeation Effects and Diurnal Emissions, Using E20 Fuels on Aging Enhanced Evaporative Emissions Certified Vehicles. Report No. E-77-2c. http://www.crcao.org/reports/recentstudies2011/E-77-2c/E-77-2c%20Final%20Report%20for%20sure%201-28-11.pdf.
- ⁵⁸ USEPA (2002). Technical Description of the Toxics Module for MOBILE6.2 and Guidance on Its Use for Emission Inventory Preparation. EPA420-R-02-011. U.S. Environmental Protection Agency; Office of Transportation and Air Quality: Ann Arbor, MI. Available at http://www.epa.gov/otaq/m6.htm.

⁴⁶ Schauer, J.J., M.J. Kleeman, G.R. Cass, and B.R.T. Simoneit. (1999) Measurement of Emissions from Air Pollution Sources, 2. C1-C30 Organic Compounds from Medium Duty Diesel Trucks. Environmental Science and Technology, vol. 33, no. 10, pp. 1578-1587, 1999.

- ⁶⁰ USEPA (2020). *Exhaust Emission Rates for Light-Duty Onroad Vehicles in MOVES3*. EPA-420-R-20-019. Office of Transportation and Air Quality. US Environmental Protection Agency. Ann Arbor, MI. November 2020. https://www.epa.gov/moves/moves-technical-reports.
- ⁶¹ USEPA (2020). *Exhaust Emission Rates of Heavy-Duty Onroad Vehicles in MOVES3*. EPA-420-R-20-018. Office of Transportation and Air Quality. US Environmental Protection Agency. Ann Arbor, MI. November 2020. https://www.epa.gov/moves/moves-technical-reports.
- ⁶² Zielinska, B., et al. (2008). Detailed Characterization and Profiles of Crankcase and Diesel Particulate Matter Exhaust Emissions Using Speciated Organics. *Environ Sci Technol*, 42 (15), 5661-5666. DOI: 10.1021/es703065h.
- ⁶³ USEPA. LA92 "Unified" Dynamometer Driving Schedule. http://www.epa.gov/otaq/standards/light-duty/la92.htm
- ⁶⁴ Okamoto et al. 2006. Unregulated Emissions from Compressed Natural Gas (CNG) Transit Buses Configured with and without Oxidation Catalyst. Environ. Sci. Technol. Vol. 40, 332-341 (value obtained from page 338, Table 6)

⁵⁹Haskew, H. M., Liberty, T. F., and McClement, D. (2004) Fuel Permeation from Automotive Systems. Prepared for the Coordinating Research Council by Harold Haskew and Associates and Automotive Testing Laboratories, Inc. September 2004. CRC Project No. E-65. http://www.crcao.org.